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(71) Applicant (for all designated States except AU BB CA GB IE LK MN MW NZ SD US): UNILEVER N.V. [NL/NL]; Weena 455, NL-3013 AL Rotterdam (NL).

(71) Applicant (for AU BB CA GB IE LK MN MW NZ SD only): UNILEVER PLC [GB/GB]; Unilever House, Blackfriars, London EC4 4BQ (GB).

(72) Inventors; and

(75) Inventors/Applicants (for US only): DAVIS, Paul, James [GB/GB]; The Hawthoms, Pavenham Road, Felmersham, Bedfordshire MK43 7EX (GB). VAN DER LOGT, Cornelis, Paul, Erik [NL/GB]; 1 Bluebell Rise (Peverel Manor Estate), Rushden, Northamptonshire MK43 7EX (GB). VERHOEUEN, Martine, Elisa [BE/GB]; 1 Tintagel Close (Manor Farm Estate), Rushden, Northamptonshire NN10 OTU (GB). WILSON, Steve [GB/GB]; 3 Aldenham Close (Goldfington), Bedford MK41 0FQ (GB).

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(54) Title: A BIFUNCTIONAL OR BIVALENT ANTIBODY FRAGMENT ANALOGUE

(57) Abstract

The invention relates to a bispecific or bivalent antibody fragment analogue comprising a binding complex containing two polypeptide chains, whereby one polypeptide chain comprises two heavy chain variable domains (VH) in series and the other polypeptide chain comprises two light chain variable domains (VL) in series, the binding complex further containing two pairs of variable domains (VH-A//VL-A and V_H-B//V_L-B). The two V_H's and/or the two V_L's are connected directly or via an intermediate peptide linker. Also a production method for such antibody fragment analogues is disclosed.

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Title: A bifunctional or bivalent antibody fragment analogue

The invention relates to new bispecific or bivalent

5 antibody fragment analogues, a process for preparing such
antibody fragment analogues and various uses of such
antibody fragment analogues.

Background of the invention and prior art

10 <u>1. Antibody structure</u>

Antibody molecules typically are Y-shaped molecules whose basic unit consist of four polypeptides, two identical heavy chains and two identical light chains, which are covalently linked together by disulfide bonds. Each of

- these chains is folded in discrete domains. The C-terminal regions of both heavy and light chains are conserved in sequence and are called the constant regions, also known as C-domains. The N-terminal regions, also known as V-domains, are variable in sequence and are responsible for the
- antibody specificity. The antibody specifically recognizes and binds to an antigen mainly through six short complementarity-determining regions located in their V-domains (see Figure 1).
- 25 In this specification abbreviations are used having the following meaning.

C-domain: Constant domain V-domain: Variable domain

 V_L : Variable domain of the light chain 30 V_B : Variable domain of the heavy chain

Fv : dual chain antibody fragment containing both a

 V_H and a V_L

scFv: single-chain Fv (V_H and V_L genetically linked

either directly or via a peptide linker)

35 CDR : Complementarity-determining region

ELISA : Enzyme Linked Immuno Sorbent Assay

PCR : Polymerase Chain Reaction

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IPTG: IsoPropyl-&-ThioGalactopyranoside

PBS : Phosphate Buffered Saline

PBST : Phosphate Buffered Saline with 0.15% Tween

TMB: 3,3',5,5'-TetraMethylBenzidine

5

It is generally known that proteolytic digestion of an antibody with papain yields three fragments. The fragment containing the CH₂ and CH₃ domains of the two heavy chains connected by the complete hinge (see Figure 1) crystallises very easily and was therefore called Fc fragment. The two other fragments are identical and were called Fab fragments, as they contained the antigen-binding site. Digestion with pepsin is such that the two Fab's remain connected via the hinge, forming only two fragments: Fc' and Fab₂.

The Fv is the smallest unit of an antibody which still contains the complete binding site (see Figure 1) and full antigen binding activity. It consists of only the V-domains of the heavy and light chains thus forming a small,

- 20 heterodimeric variable fragment or Fv. Fv's have a molecular weight of about 25 kD, which is only one sixth of the parent whole antibody (in the case of an IgG).
 Previously Fv's were only available by proteolysis in a select number of cases (Givol, 1991). The production of
- 25 Fv's can now be achieved more routinely using genetic engineering methods through cloning and expressing DNA encoding only the V-domains of the antibody of interest.

 Smaller fragments, such as individual V-domains (Domain Antibodies or dABs, Ward et al., 1989), and even individual
- 30 CDR's (Williams et al., 1989; Taub et al., 1989) were shown to retain the binding characteristics of the parent antibody. However, this is not achievable on a routine basis: most naturally occurring antibodies need both a $V_{\rm H}$ and a $V_{\rm L}$ to retain full immunoreactivity. For example, in
- 35 the case of V_H D1.3 (Ward et al., 1989), although it still binds hen egg lysozyme (HEL) with an affinity close to that of the parent antibody, it was shown that loss of

specificity was observed in that it can no longer distinguish turkey lysozyme from HEL, whereas the Fv can (Berry and Davies, 1992). Although murine dABs can be obtained more routinely from spleen libraries (Ward et al., 5 1989), the approach is unsustainable because of the many problems associated with their production and physical behaviour: expression is extremely poor, affinity tends to be low, stability and solubility in water is low, and nonspecific binding is usually very high. According to the 10 literature a possible explanation of these undesirable characteristics is the exposure of the hydrophobic residues which are normally buried in the V_H-V_t interface. The exposed hydrophobic patches are thought to contribute to aggregation of the protein inside the cells and/or in the 15 culture medium, leading to poor expression and/or poor solubility (Anthony et al., 1992; Ward et al., 1989). The hydrophobic patches can also explain the high non-specific binding described by Berry and Davies, 1992. These problems clearly limit the usefulness of these molecules. 20 Most of the Camelid antibodies appear to be an exception to this rule in that they only need one V-domain, namely V_H , to specifically and effectively bind an antigen (Hamers-

Castermans et al., 1993). In addition, preliminary data indicate that they seem not to suffer from the

25 disadvantages of mouse dABs, as these camelid antibodies or fragments thereof are soluble and have been shown to express well in yeast and Aspergillus moulds. These observations can have important consequences for the production and exploitation of antibody-based products, see

patent application WO 94/25591 (UNILEVER et al., first priority date 29.04.93).

2. Production of antibody fragments

Several microbial expression systems have already been developed for producing active antibody fragments, e.g. the production of Fab in various hosts, such as E. coli (Better et al., 1988, Skerra and Plückthun, 1988, Carter et al.,

1992), yeast (Horwitz et al., 1988), and the filamentous fungus Trichoderma reesei (Nyyssönen et al., 1993) has been described. The recombinant protein yields in these alternative systems can be relatively high (1-2 g/l for Fab 5 secreted to the periplasmic space of E. coli in high cell density fermentation, see Carter et al., 1992), or at a lower level, e.g. about 0.1 mg/l for Fab in yeast in fermenters (Horwitz et al., 1988), and 150 mg/l for a fusion protein CBHI-Fab and 1 mg/l for Fab in Trichoderma 10 in fermenters (Nyyssonen et al., 1993) and such production is very cheap compared to whole antibody production in mammalian cells (hybridoma, myeloma, CHO). Although the latter can give yields of the order of 1 g/l in high cell density fermentation, it is a time-consuming and very 15 expensive manufacturing method resulting in a cost price of about 1000 £/gram of antibody. It was further demonstrated that plants can be used as hosts for the production of both whole antibodies (Hiatt et al., 1989) and scFv's (Owen et al., 1992, Firek et al., 1993), whereby yields of upto 0.5% 20 of the total soluble protein content in tobacco leaves were mentioned.

The fragments can be produced as Fab's or as Fv's, but additionally it has been shown that a V_H and a V_L can be genetically linked in either order by a flexible polypeptide linker, which combination is known as an scFv (Bird et al. (1988), Huston et al. (1988), and granted patent EP-B-0281604 (GENEX/ENZON LABS INC.; first priority date 02-09-1986).

30 3. Bivalent and bispecific antibodies and antibody ____fragments

The antibody fragments Fab, Fv and scFv differ from whole antibodies in that the antibody fragments carry only a single antigen-binding site. Recombinant fragments with two binding sites have been made in several ways, for example, by chemical cross-linking of cysteine residues introduced at the C-terminus of the V_H of an Fv (Cumber et al., 1992),

or at the C-terminus of the V_L of an scFv (Pack and Plückthun, 1992), or through the hinge cysteine residues of Fab's (Carter et al., 1992). Another approach to produce bivalent antibody fragments is described by Kostelny et al. (1992) and Pack and Plückthun (1992) and is based on the inclusion of a C-terminal peptide that promotes dimerization.

When two different specificities are desired, one can generate bispecific antibody fragments. The traditional approach to generate bispecific whole antibodies was to fuse two hybridoma cell lines each producing an antibody having the desired specificity. Because of the random association of immunoglobulin heavy and light chains, these hybrid hybridomas produce a mixture of up to 10 different

- heavy and light chain combinations, only one of which is the bispecific antibody (Milstein and Cuello, 1983). Therefore, these bispecific antibodies have to be purified with cumbersome procedures, which considerably decrease the yield of the desired product.
- Alternative approaches include in-vitro linking of two antigen specificities by chemical cross-linking of cysteine residues either in the hinge or via a genetically introduced C-terminal Cys as described above. An improvement of such in vitro assembly was achieved by using recombinant fusions of Fab's with peptides that promote formation of heterodimers (Kostelny et al., 1992). However, the yield of bispecific product in these methods is far less than 100%.

A more efficient approach to produce bivalent or bispecific antibody fragments, not involving in vitro chemical assembly steps, was described by Holliger et al. (1993). This approach takes advantage of the observation that scFv's secreted from bacteria are often present as both monomers and dimers. This observation suggested that the $V_{\rm H}$ and $V_{\rm L}$ of different chains can pair, thus forming dimers and larger complexes. The dimeric antibody fragments, also named "diabodies" by Hollinger et al., in fact are small

bivalent antibody fragments that assembled in vivo. By linking the V_H and V_L of two different antibodies 1 and 2, to form "cross-over" chains $V_H 1 V_L 2$ and $V_H 2 - V_L 1$ (see Figure 2B), the dimerisation process was shown to reassemble both 5 antigen-binding sites. The affinity of the two binding sites was shown to be equal to the starting scFv's, or even to be 10-fold increased when the polypeptide linker covalently linking V_H and V_L was removed, thus generating two proteins each consisting of a VH directly and 10 covalently linked to a V_L not pairing with the V_H (see Figure 2C). This strategy of producing bispecific antibody fragments was also described in several patent applications. Patent application WO 94/09131 (SCOTGEN LTD; priority date 15.10.92) relates to a bispecific binding 15 protein in which the binding domains are derived from both a V_H and a V_L region either present at two chains or linked in an scFv, whereas other fused antibody domains, e.g. Cterminal constant domains, are used to stabilise the dimeric constructs. Patent application WO 94/13804 20 (CAMBRIDGE ANTIBODY TECHNOLOGY / MEDICAL RESEARCH COUNCIL; first priority date 04.12.92) relates to a polypeptide containing a V_H and a V_L which are incapable of associating with each other, whereby the V-domains can be connected with or without a linker. Mallender and Voss, 1994 (also described in patent application WO 94/13806; DOW CHEMICAL CO; priority date

Mallender and Voss, 1994 (also described in patent application WO 94/13806; DOW CHEMICAL CO; priority date 11.12.92) reported the in vivo production of a single-chain bispecific antibody fragment in *E. coli*. The bispecificity of the bivalent protein was based on two previously produced monovalent scFv molecules possessing distinct specificities, being linked together at the genetic level by a flexible polypeptide linker. The thus formed V_H1-linker-V_L1-linker-V_H2-linker-V_L2 fragment (see Figure 2A) was shown to contain both antigen binding specificities 1 and 2. (1= anti-fluorescein, 2= anti-single-stranded DNA).

Traditionally, whenever single-chain antibody fragments are referred to, a single molecule consisting of one heavy chain linked to one (corresponding) light chain in the presence or absence of a polypeptide linker is implicated.

- When making bivalent or bispecific antibody fragments through the 'diabody' approach (Holliger et al., (1993) and patent application WO 94/09131) or by the 'double scFv' approach (Mallender and Voss, 1994 and patent application WO 94/13806), again the $V_{\rm H}$ is linked to a (the
- 10 corresponding) V_L .

It is realised that claims 32 and 33 of patent application WO 93/11161 (ENZON INC.; priority date 25.11.91) and the corresponding passages in that specification on page 22, lines 1-10 may read on a polypeptide comprising two $V_L{}^\prime s$

- 15 fused together via a flexible polypeptide linker, and on a polypeptide comprising two V_{H} 's fused together via a flexible polypeptide linker, respectively. However, no examples were given to substantiate this approach, thus it was in fact a hypothetical possibility instead of an
- 20 actually produced compound.

A skilled person would not have expected that such approach would be viable for at least three reasons. Firstly, it is widely recognised that immunoglobulin heavy chains (excluding the above described camel immunoglobulins) have

- very limited solubility and spontaneously precipitate out of aqueous solution when isolated from their light chain partners. Secondly, several groups have shown (Ward et al., 1989, Berry and Davies, 1992, and Anthony et al., 1992) that expression of V_H's in the absence of V_L's is hampered
- by extremely poor yields of unstable product with many undesirable properties, e.g. non-specific binding. Thirdly in patent application WO 94/13804 it was described on page 31 lines 10-12, that in computer modelling experiments they could not model as heterodimers V_H-V_H and V_L-V_L given the
- onstraints of short linkers.

 Thus the simple suggestion given in patent application WO
 93/11161 is not an enabling disclosure leading a skilled

person to try with a reasonable expectation of success whether such suggestion would work; therefore, that patent application should not be considered as relevant prior art for the present invention.

5

Summary of the invention

The present invention provides a **bispecific** or **bivalent** antibody fragment analogue, which comprises a binding complex containing two polypeptide chains, one of which comprises two times a variable domain of a heavy chain (V_H) in series and the other comprises two times a variable domain of a light chain (V_L) in series.

In one aspect of the invention one chain of the antibody fragment analogue comprises a first V_H (V_H-A) connected to a second V_H (V_H-B) and the other chain comprises a first V_L (V_L-A) connected to a second V_L (V_L-B). In a preferred embodiment of this aspect one chain comprises a first V_H (V_H-A) followed by a second V_H (V_H-B), thus [V_H-A * V_H-B], and the other chain comprises a first V_L (V_L-A) preceded by a second V_L (V_L-B), thus [V_L-B * V_L-A]. For some embodiments of this aspect the two V_H's are directly connected to each other, but for other embodiments of this aspect of the invention the two V_L's are directly connected to each other. According to another embodiment of this aspect of the invention the two V_H's are connected to each other by a linker and also the two V_L's are connected to each other by

According to a special embodiment of this aspect of the invention one chain comprises a first V_H (V_{H} -A) followed by a second V_H (V_{H} -B), thus [V_{H} -A * V_{H} -B], and the other chain comprises a first V_L (V_L -A) followed by a second V_L (V_L -B), thus [V_L -A * V_L -B], and in which the two V_H 's are connected to each other by a linker and also the two V_L 's are connected to each other by a linker, whereas each linker

comprises at least 10 amino acid residues.

a linker. Such a linker usually comprises at least one

amino acid residue.

According to the above aspect of the invention with A being different from B there are provided **bispecific** antibody fragment analogues.

According to another aspect of the invention the

5 specificities A and B are the same resulting in a bivalent antibody fragment.

According to a further aspect of the invention the bispecific or bivalent antibody fragment analogues can be used in a diagnostic technique or for immunoassays, in a

- 10 purification method, for therapy, or in other methods in which immunoglobulins or fragments thereof are used. Such uses are well-known in the art.
 - The invention also provides a process for producing the antibody fragments of the invention in that a host is
- transformed by incorporating into that host a DNA encoding the two V_{H} 's with or without a connecting linker and a DNA encoding the two V_{L} 's with or without a connecting linker. Preferably the two DNA's are placed in a dicistronic arrangement.
- It is also possible that the two linked V_H 's and the two linked V_L 's are produced separately by different hosts, after which the linked V_H 's produced by one host can be combined with the linked V_L 's produced by the other host. The hosts can be selected from the group consisting of
- prokaryotic bacteria of which examples are Gram-negative bacteria, e.g. *B. coli*, and Gram-positive bacteria, e.g. *B. subtilis* or lactic acid bacteria, lower eukaryotes examples of which are yeasts, e.g. belonging to the genera Saccharomyces, Kluyveromyces, or Trichoderma, moulds, e.g.
- belonging to the genera Aspergillus and Neurospora, and higher eukaryotes, examples of which are plants, e.g. tobacco, and animal cells, examples of which are myeloma cells and CHO cells. The techniques to transform a host by genetic engineering methods in order to have a desirable
- polypeptide produced by such host are well-known to persons skilled in the art as is evident from the literature

mentioned above under the heading "Background of the invention and prior art".

Brief description of the drawings 5 Figure 1 depicts in schematic form the structure of a typical antibody (immunoglobulin) molecule. shows a schematic representation of published Figure 2 arrangements of heavy and light chain V-domain gene fragments that have been proven to 10 produce bispecific antibody fragments. Figure 3 shows in diagrammatic form the suggested arrangement of the V-domains of a double head antibody fragment according to the invention with the V-domains in the following order: 15 $V_{\mu}A - V_{\mu}B + V_{\mu}B - V_{\mu}A$. Figure 4 shows the nucleotide sequence of the EcoRI-HindIII insert of pUR.4124 containing DNA (see SEQ ID NO: 1) encoding V_LLys-Linker-V_HLys (see SEO ID NO: 2). 20 Figure 5 shows the nucleotide sequence of the HindIII-EcoRI insert of plasmid Fv.3418 (see SEQ ID NO: 3) containing DNA encoding pelB leader-V_H3418 (see SEQ ID NO: 4) and DNA encoding pelB leader-V_L3418 (see SEQ ID NO: 5). 25 Figure 6 shows the nucleotide sequence of the HindIII-EcoRI insert of plasmid Fv.4715-myc (see SEQ ID NO: 6) containing DNA encoding pelB leader-V_H4715 (see SEQ ID NO: 7) and DNA encoding pelB leader-V_L4715-Myc tag (see SEQ ID NO: 8)... 30 Figure 7 shows the nucleotide sequence of the HindIII-BcoRI insert of scFv.4715-myc containing DNA (see SEQ ID NO: 9) encoding pelB leader-V_H4715-Linker-V_L4715-Myc tag (see SEQ ID NO: 10).

Figure 8 a/b shows the nucleotide sequence of the HindIII-

EcoRI insert of pGOSA.E (see SEQ ID NO: 11) containing DNA encoding pelB leader-V_H4715-

Linker-V,3418 (see SEQ ID NO: 12) and DNA encoding pelB leader-VL3418-Linker-VR4715 (see SEQ ID NO: 13). Figure 9 gives an overview of the oligonucleotides and 5 their positions in pGOSA.E that can be used to replace V-domain gene fragments. Figure 10 illustrates the amino acid sequence of the Vu- V_H and V_L-V_L domain junctions in fusion polypeptides GOSA.E (see amino acids 114-145 10 in SEQ ID NO: 12 and amino acids 102-128 in SEQ ID NO: 13), GOSA.V (see SEQ ID NO: 30 and amino acids 102-128 in SEQ ID NO: 13), GOSA.S (see amino acids 114-145 in SEQ ID NO: 12 and SEQ ID NO: 31) and GOSA.T (see SEQ ID NO: 30 15 and SEQ ID NO: 31). Figure 11 shows the specificity of Streptococcus binding of scFv.4715-myc. Figure 12 shows the specificity of glucose oxidase targeting onto the surface of various 20 Streptococcus strains by GOSA.E. Figure 13 shows the specificity of glucose oxidase targeting onto the surface of various Streptococcus strains by GOSA.V. Figure 14 shows the specificity of glucose oxidase 25 targeting onto the surface of various Streptococcus strains by GOSA.S. Figure 15 shows the specificity of glucose oxidase targeting onto the surface of various Streptococcus strains by GOSA.T. 30 Figure 16 shows the results of an ELISA. Individual fractions of a gelfiltration experiment using partially purified GOSA.E as feedstock were tested for glucose oxidase and Streptococcus sanguis bispecific binding activity. 35 Figure 17 shows the results of an ELISA. Individual fractions of a gelfiltration experiment using partially purified GOSA.V as feedstock were

tested for glucose oxidase and Streptococcus sanguis bispecific binding activity. Figure 18 shows the results of an ELISA. Individual fractions of a gelfiltration experiment using 5. partially purified GOSA.S as feedstock were tested for glucose oxidase and Streptococcus sanguis bispecific binding activity. Figure 19 shows the results of an ELISA. Individual fractions of a gelfiltration experiment using 10 partially purified GOSA.T as feedstock were tested for glucose oxidase and Streptococcus sanguis bispecific binding activity. shows the source of fragment PCR.I BstEII/SacI Figure 20 Figure 21 shows the source of fragment PCR.II Sfil/EcoRI 15 Figure 22 shows the source of fragment PCR.III NheI/SacI Figure 23 shows the source of fragment PCR.IV XhoI/EcoRI Figure 24 shows the source of fragment PCR.V Sall/EcoRI Figure 25 shows the source of fragment PCR.VI Sfil/NheI Figure 26 shows the source of fragment PCR.VII 20 BstEII/NheI Figure 27 shows the source of fragment PCR.VIII XhoI/EcoRI Figure 28 shows the source of fragment PCR.IX BstEII/NheI Figure 29 25 shows the source of fragment PCR.X PstI/EcoRI shows the construction of plasmid pGOSA.A Figure 30 Figure 31 shows the construction of plasmid pGOSA.B Figure 32 shows the construction of plasmid pGOSA.C Figure 33 shows the construction of plasmid pGOSA.D 30 Figure 34 shows the construction of plasmid pGOSA.E Figure 35 shows the construction of plasmid pGOSA.V Figure 36 shows the construction of plasmid pGOSA.S Figure 37 shows the construction of plasmid pGOSA.T Figure 38 a/b shows the construction of plasmid pGOSA.G 35 Figure 39 shows the construction of plasmid pGOSA.J Figure 40 shows the construction of plasmid pGOSA.Z shows the construction of plasmid pGOSA.AA Figure 41

Figure	42	shows	the	construction	of	plasmid	pGOSA.AB
Figure	43	shows	the	construction	of	plasmid	pGOSA.L
Figure	44	shows	the	${\tt construction}$	of	plasmid	pGOSA.Y
Figure	45 .	shows	the	construction	of	plasmid	pGOSA.X
Figure	46	shows	the	construction	of	plasmid	pGOSA.AC
Figure	47	shows	the	construction	of	plasmid	pGOSA.AD.

- Table 1 shows the nucleotide sequence of the oligonucleotides used to produce the constructs described in this specification. Restriction sites encoded by these primers are underlined.
 - Table 2 gives an overview of all GOSA constructs described in this specification.
- Table 2A describes intermediate constructs that were not further tested.

 Table 2B describes the dicistronic constructs.

 Table 2C describes the monocistronic constructs.

20 <u>Detailed description of the invention</u>

In this specification the construction of an antibody fragment analogue consisting of a two chain protein complex is described, in which one of the chains consists of two heavy chain V-domains and the other chain consists of the two corresponding light chain V-domains in either order. The variable domains are linked either directly or through a polypeptide linker. Subsequent molecular modelling of this combination suggested that the protein chains could fold such that both binding sites are fully accessible, provided that the connecting linkers are kept long enough to span 30 to 35 Å.

Whereas in patent application WO 93/11161 it is explicitly described that for the above described bispecific complexes two flexible polypeptide linkers in the self assembling complex are required, the present invention illustrated here describes in particular the construction of a two chain protein complex containing only one linker or no

linkers at all. The latter antibody fragment analogue thus consists of a two chain protein complex containing one polypeptide chain comprising heavy chain V-domains fused directly together and another polypeptide chain comprising the corresponding light chain V-domains fused together, both fusions in the absence of linkers. But also two chain protein complexes in which each chain comprises a linker between the two variable domains can be used as antibody fragment analogues according to the invention as described below with construct pGOSA.E. However, the two chain complexes containing only one linker or no linker at all are preferred. The abbreviation GOSA used in this specification relates to a combination of glucose oxidase and Streptococcus sanguis.

15

In this specification evidence is provided that these antibody fragment analogues ("double heads") contain both antigen binding specificities of the Fv's used to generate these bispecific antibody fragments. It is exemplified that these type of constructs according to the invention can be used to target the enzyme glucose oxidase to whole bacteria, using antibody fragments derived from hybridomas expressing antibodies directed against these antigens.

The present invention is now described by reference to some specific examples, which are included for purposes of illustration only and are not intended to limit the scope of the invention.

30

EXAMPLES

General experimental

Strains, Plasmids and Media

All cloning steps were performed in E. coli JM109 (endA1, recA1, gyrA96, thi, hsdR17(r_K, m_K⁺), relA1, supB44, Δ (lacproAB), [F', traD36, proAB, lacI^QZ Δ M15]. E. coli cultures were grown in 2xTY medium (16 g tryptone, 10 g yeast extract, 5 g NaCl per litre H₂O), where indicated

supplemented with 2% glucose and/or 100 µg/ml ampicillin. Transformations were plated out on SOBAG plates (20 g tryptone, 5 g yeast extract, 15 g agar, 0.5 g NaCl per litre H₂O plus 10 mM MgCl₂, 2% glucose, 100 µg/ml ampicillin) The expression vectors used are derivatives of pUC19. The oligonucleotide primers used in the PCR reactions were synthesized on an Applied Biosystems 381A DNA Synthesiser by the phosphoramidite method.

10 Expression of GOSA constructs

Colonies from freshly transformed JM109 plated onto SOBAG plates were used to inoculate 2xTY medium supplemented with 100 μ g/ml ampicillin, 2% glucose. Cultures were shaken at 37°C to an OD₆₀₀ in the range of 0.5 to 1.0. Cells were

- pelleted by centrifugation and the supernatant was removed. The pelleted cells were resuspended in 2xTY medium with 100 μ g/ml ampicillin, 1 mM IPTG, and grown for a further 18 hours at 25°C. Cells were pelleted by centrifugation and the supernatant, containing the secreted chains, used
- directly in an ELISA. The proteins in the periplasm of the pelleted cells were extracted by resuspending the cell pellet in 1/20 of the original culture volume of lysis buffer (20% sucrose, 200 mM Tris-HCl pH 7.5, 1 mM EDTA, 500 μg/ml lysozyme). After incubation at 25°C for 20 minutes an
- equal volume of H₂O was added and the incubation was continued for another 20 minutes. The suspension was spun at 10.000 g for 15 minutes and the supernatant containing the periplasmic proteins was used directly in an ELISA.

30 ELISA

96 well ELISA plates (Greiner HC plates) were activated overnight at 37°C with 200 μ l/well of an 1/10 dilution of an over night culture of *Streptococcus* cells in 0.05 M sodium carbonate buffer at pH=9.5. Following one wash with PBST, the antigen sensitised plates were pre-blocked for 1 hour at 37°C with 200 μ l/well blocking buffer (2% BSA, 0.15% Tween in PBS). Samples containing 50 μ l blocking

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buffer plus 50 μl culture supernatants or periplasmic cell extracts (neat or diluted with PBS) were added to the Streptococcus sensitised plate and incubated for 2 hours at 37°C. Following 4 washes with PBS-T, 100 μl of blocking buffer containing glucose oxidase (50 μg/ml) was added to every well. After incubation at 37°C for 1 hour unbound glucose oxidase was removed by 4 washes with PBS-T. Bound glucose oxidase was detected by adding 100 μl substrate to each well (70 mM Na-citrate, 320 mM Na-phosphate, 27 mg/ml glucose, 0.5 μg/ml HRP, 100 μg/ml TMB). The colour reaction was stopped after 1 hour by the addition of 35 μl 2 M HCl and the A450 was measured (compare Figures 11/15).

Affinity purification of GOSA antibody fragments

- 15 GOSA.E, GOSA.V, GOSA.S and GOSA.T were partially purified by affinity chromatography. 100 ml periplasmic extract of each of these constructs was loaded onto a Glucose-oxidase-Sepharose column (CNBr-Sepharose, Pharmacia) prepared according to the manufacturer's instructions. After
- extensive washes with PBS the bound GOSA antibody fragments were eluted in 0.1M glycine buffer at pH=2.8. The fractions were neutralised with Tris and analysed by polyacrylamide gel electrophoresis followed by silver staining and tested for the presence of double head activity.

25

EXAMPLE 1. Construction of the pGOSA double head expression vectors

- In this Example the construction of a two chain protein complex is described, in which one of the chains consists of two heavy chain V-domains and the other chain consists of the two corresponding light chain V-domains. The variable domains are linked either directly or through a
- polypeptide linker. The expression vectors used are derivatives of a pUC19 derived plasmid containing a HindIII-BcoRI fragment that in the case of plasmid

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scFv.4715-myc contains a DNA fragment encoding one pelB signal sequence fused to the N-terminus of the Vu that is directly linked to the corresponding V_L of the antibody through a connecting flexible peptide linker, (Gly,Ser), 5 (present in SEQ ID NO: 2 as amino acids 109-123 and in SEQ ID NO: 10 as amino acids 121-135), thus generating a single-chain molecule (see Figure 7). In the dual-chain Fv and the pGOSA expression vectors, the DNA fragments encoding both the V_H and V_L of the antibody are preceded by a ribosome binding site and a DNA sequence encoding the pelB signal sequence in an artificial dicistronic operon under the control of a single inducible promoter (see Figures 5, 6 and 8). Expression of these constructs is driven by the inducible lacZ promoter. The 15 nucleotide sequence of the HindIII-EcoRI inserts of the plasmids pUR.4124, Fv.3418, Fv.4715-myc and scFv.4715-myc constructs used for the generation of the bispecific antibody fragments are given in Figures 4-7, respectively. Moreover, a culture of E. coli cells harbouring plasmid scFv.4715-myc and a culture of E. coli cells harbouring plasmid Fv.3418 were deposited under the Budapest Treaty at the National Collection of Type Cultures (Central Public Health Laboratory) in London (United Kingdom) with deposition numbers NCTC 12916 and NCTC 12915, respectively. In agreement with Rule 28 (4) EPC, or a similar arrangement for a State not being a Contracting State of the BPC, it is hereby requested that a sample of such deposit, when requested, will be submitted to an expert only. The construction of pGOSA.E (see Figure 8 for the HindIII-EcoRI insert of pUC19) involved several cloning steps. The appropriate restriction sites in the various domains were introduced by PCR directed mutagenesis using the oligonucleotides listed in Table 1 below. The pGOSA.E derivatives pGOSA.V, pGOSA.S and pGOSA.T with only one or no linker sequence are derived from the pGOSA.E construct by removing the linker sequences by means of PCR directed

mutagenesis with oligonucleotides listed in Table 1 below.

Table 1.

	DBL.1	5'-CAC CAT CTC CAG AGA CAA TGG CAA	G-3'			
			(=SEQ ID NO: 14			
	DBL.2	5'-GAG CGC GAG CTC GGC CGA ACC GGC CIGA TCC GCC				
5		ACC GCC AGA GCC-3'	(=SEQ ID NO: 15			
	DBL.3	5'-CAG GAT CCG GCC GGT TCG GCCI CAG	GTC CAG CTG			
		CAA CAG TCA GGA-3'	(=SEQ ID NO: 16			
	DBL.4	5'-CTA CAT GAA TTC ² GCT AGC ³ TTA TTA	TGA GGA GAC			
		GGT GAC GGT GGT CCC TTG GC-3	(=SEQ ID NO: 17			
10	DBL.5	5'-TAA TAA GCT AGC3 GGA GCT GCA TGC	AAA TTC TAT			
		TTC-3'	(= SEQ ID NO: 18			
	DBL.6	5'-ACC AAG CTC GAG4 ATC AAA CGG GG-3	3'(= SEQ ID NO: 19			
	DBL.7	5'-AAT GTC GAA TTC' GTC GAC' TCC GCC	ACC GCC AGA			
		GCC-3	(= SEQ ID NO: 20			
15	DBL.8	5'-ATT GGA GTC GAC' ATC GAA CTC ACT	CAG TCT CCA			
		TTC TCC-3'	(= SEQ ID NO: 21			
	DBL.9	5'-TGA AGT GAA TTC' GCG GCC GC"T TAT	TAC CGT TTG			
		ATT TCG AGC TTG GTC CC-3'	(= SEQ ID NO: 22			
	DBL.10	5'-CGA ATT CGG TCA CC8G TCT CCT CAC	AGG TCC AGT			
20		TGC AAC AG-3'	(= SEQ ID NO: 23			
	DBL.11	5'-CGA ATT CTC GAG4 ATC AAA CGG GAC	ATC GAA CTC			
		ACT CAG TCT CC-3'	(= SEQ ID NO: 24			
	DBL.12	5'-CGA ATT CGG TCA CC G TCT CCT CAC	AGG TGC AGT			
		TGC AGG AG-3'	(= SEQ ID NO: 25			
25	PCR.51	5'-AGG T(C/G)(A/C) A(C/A)C TGC AG ⁷ (C/G)	AGT C(A/T)G			
		G-3'	(= SEQ ID NO: 26			
	PCR.89	5'-TGA GGA GAC GGT GAC C'SGT GGT CCC	C TTG GCC CC-3'			
			(= SEQ ID NO: 27			
	PCR.90	5'-GAC ATT GAG CTC' ACC CAG TCT CCA-	3(= SEQ ID NO: 28			
30	PCR.116	5'-GTT AGA TCT CGA G'CT TGG TCC C-3'	(= SEQ ID NO: 29			
	l = SfiI, 2	= $EcoRI$, $3 = NheI$, $4 = XhoI$, $5 = SalI$, $6 = NotI$, $7 = Pst$	I, 8=BstEII, 9=SacI			

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These three constructs lack some of the restriction sites at the new joining points. The V_HA-V_HB gene fragment without a linker lacks the 5' V_HB SfiI site. The V_LB-V_LA gene fragment without a linker lacks the 5' V_LA SalI site.

The position of the oligonucleotides in the pGOSA constructs given in Table 1 are shown in Figure 9. The pGOSA expression vectors and the oligonucleotides in Table 1 have been designed to enable most specificities to be cloned into the pGOSA constructs. Figure 10 shows the amino acid sequence of the junctions between the V_HA-V_HB and V_LB-V_LA fragments encoded by DNA present in pGOSA.E, pGOSA.V, pGOSA.S and pGOSA.T. A more detailed description of the preparation of pGOSA.E, pGOSA.V, pGOSA.S and pGOSA.T is given in Example 5.

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EXAMPLE 2. Bifunctional binding activity of GOSA double heads

In this Example we provide evidence that the above described molecules ("double heads"), i.e. the two chain protein complexes, contain both antigen binding specificities of the Fv's used to generate these multifunctional antibody fragment analogues. Figure 12-15 show that GOSA.E, GOSA.V, GOSA.S and GOSA.T can be used to specifically target the enzyme glucose oxidase to several Streptococcus sanguis strains using antibody fragments derived from hybridoma's expressing antibodies directed against these antigens.

Comparison of the binding specificity of the GOSA constructs (see Figures 12-15) and the binding specificity of the scFv.4715-myc (see Figure 11) shows that the fine specificity of the anti-Streptococcus sanguis scFv.4715 is preserved in the GOSA "double heads".

EXAMPLE 3. FPLC analysis of GOSA double heads

Partially purified GOSA.E, GOSA.V, GOSA.S and GOSA.T samples (estimated to be 50-80% pure by polyacrylamide qel 5 electrophoresis) were analysed on a Pharmacia FPLC Superose 12 column. The analysis was performed using PBS at a flow rate of 0.3 ml/minute. Eluate was monitored at 280 nm and 0.3 ml fractions were collected and analysed by ELISA. Usually GOSA.E, GOSA.V, GOSA.S and GOSA.T samples only gave 10 one GOSA double head activity peak as determined by ELISA (see Figures 16-19). The position of this peak in the elution pattern indicated that the molecular weight of the GOSA double head is 40-50 kD. Since this molecular weight corresponds to the expected molecular weight of the $V_{\rm H}2$ + 15 V₁2 double head dimer, the conclusion is justified that GOSA.E, GOSA.V, GOSA.S and GOSA.T are primarily produced as dimeric molecules. Occasionally an activity peak with an apparent molecular weight of ~200 kD was observed (see Figure 16). The presence of Glucose Oxidase activity in 20 these fractions (data not shown) indicate that these fractions contain GOSA double head complexed with glucose oxidase that was eluted with the GOSA sample from the glucose oxidase-sepharose affinity matrix.

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EXAMPLE 4. Production of other double heads

anti-hen-egg-lysozyme /anti-glucose oxidase,

anti-huIgG / anti-glucose oxidase,

The methods described in the previous Examples were used to produce other double heads, which also appeared to be

30 active against the antigens for which they were developed. These other double heads had the following specificities: anti-S. sanguis / anti-beta-HCG, anti-S. sanguis / anti-urease, anti-S. sanguis / anti-hen-egg-lysozyme,

35 anti-beta-HCG / anti-hen-egg-lysozyme,

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anti-urease / anti-glucose oxidase, anti-lacto-peroxidase / anti-glucose oxidase, anti-alpha-HCG / anti-glucose oxidase, and anti-reactive-Red-6 / anti-glucose oxidase.

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EXAMPLE 5. Detailed description of the preparation of intermediate constructs pGOSA.A, pGOSA.B pGOSA.C and pGOSA.D and their use for the preparation of plasmid pGOSA.E and its derivatives pGOSA.V, pGOSA.S and pGOSA.T

Oligonucleotides and PCR

The primary structures of the oligonucleotide primers used in the construction of the bispecific 'pGOSA' constructs are shown in Table 1 above. Reaction mixture used for amplification of DNA fragments were 10 mM Tris-HCl, pH 8.3, 2.5 mM MgCl₂, 50 mM KCl, 0.01% gelatin (w/v), 0.1% Triton X-100, 400 mM of each dNTP, 5.0 units of Vent DNA polymerase (New England Biolabs), 100 ng of template DNA, and 500 ng of each primer (for 100 µl reactions). Reaction conditions were: 94°C for 4 minutes, followed by 33 cycles of each 1 minute at 94°C, 1 minute at 55°C, and 1 minute 72°C.

25 Plasmid DNA\Vector\Insert preparation and ligation\transformation.

Plasmid DNA was prepared using the 'Qiagen P-100 Midi-DNA Preparation' system. Vectors and inserts were prepared by digestion of 10 µg (for vector preparation) or 20 µg (for insert preparation) with the specified restriction endonucleases under appropriate conditions (buffers and temperatures as specified by suppliers). Modification of the DNA ends with Klenow DNA polymerase and dephosphorylation with Calf Intestine Phosphorylase were performed according to the manufacturers instructions. Vector DNAs and inserts were separated through agarose gel electrophoresis and purified with DEAE-membranes NA45

(Schleicher & Schuell) as described by Maniatis et al.

Ligations were performed in 20 µl volumes containing 30 mM

Tris-HCl pH 7.8, 10 mM MgCl₂, 10 mM DTT, 1 mM ATP, 300-400

ng vector DNA, 100-200 ng insert DNA and 1 Weiss unit T₄ DNA

5 ligase. After ligation for 2-4 h at room temperature, CaCl₂

competent E. coli JM109 (Maniatis) were transformed using

7.5 µl ligation reaction. The transformation mixtures were

plated onto SOBAG plates and grown overnight at 37°C.

Correct clones were identified by restriction analysis and

10 verified by automated dideoxy sequencing (Applied

Biosystems).

Restriction digestion of PCR products

- Following amplification each reaction was checked for the presence of a band of the appropriate size by agarose gel electrophoresis. One or two 100 μ l PCR reaction mixtures of each of the PCR reactions PCR.I PCR.X (Figure 20-29), together containing approximately 2-4 μ g DNA product were subjected to phenol-chloroform extraction, chloroform
- extraction and ethanol precipitation. The DNA pellets were washed twice with 70% ethanol and allowed to dry. Next, the PCR products were digested overnight (18 h) in the presence of excess restriction enzyme in the following mixes at the specified temperatures and volumes.
- 25 PCR.I:
 - 50 mM Tris-HCl pH 8.0, 10 mM MgCl₂, 50 mM NaCl, 4 mM spermidine, 0.4 μ g/ml BSA, 4 μ l (= 40 U) SacI, 4 μ l (= 40 U) BstEII, in 100 μ l total volume at 37°C. PCR.II:
- 30 10 mM Tris-Acetate pH 7.5, 10 mM MgAc2, 50 mM KAc (1x
 "One-Phor-All" buffer ex Pharmacia), 4 μl (= 48 U) SfiI, in
 50 μl total volume at 50°C under mineral oil. After
 overnight digestion, PCR.II-SfiI was digested with EcoRI
 (overnight at 37°C) by the addition of 16 μl H2O, 30 μl 10x
 35 "One-Phor-All" buffer (Pharmacia) (100 mM Tris-Acetate pH

7.5, 100 mM MgAc₂, 500 mM KAc) and 4 μ l (= 40 U) EcoRI.

PCR.III:

10 mM Tris-Acetate pH 7.5, 10 mM MgAc₂, 50 mM KAc (1x "One-Phor-All" buffer {Pharmacia}), 4 μ l (= 40 U) NheI, 4 μ l (= 40 U) SacI, in 100 μ l total volume at 37°C.

5 PCR.IV:

20 mM Tris-Acetate pH 7.5, 20 mM MgAc₂, 100 mM KAc (2x "One-Phor-All" buffer {Pharmacia}), 4μ l (= 40 U) XhoI, 4 μ l (= 40 U) EcoRI, in 100 μ l total volume at 37°C. PCR.V:

- 10 20 mM Tris-Acetate pH 7.5, 20 mM MgAc₂, 100 mM KAc (2x "One-Phor-All" buffer {Pharmacia}), 4 μ l (= 40 U) Sall, 4 μ l (= 40 U) EcoRI, in 100 μ l total volume at 37°C. PCR.VI:
- 10 mM Tris-Acetate pH 7.5, 10 mM MgAc₂, 50 mM KAc (1x "One-Phor-All" buffer {Pharmacia}), 4 μl (= 48 U) SfiI, in 50 μl total volume at 50°C under mineral oil. After overnight digestion, PCR.VI-SfiI was digested with NheI (overnight at 37°C) by the addition of 41 μl H₂O, 5 μl 10x "One-Phor-All" buffer (Pharmacia) (100 mM Tris-Acetate pH
- 20 7.5, 100 mM MgAc₂, 500 mM KAc) and 4 μ l (= 40 U) NheI. PCR.VII:

50 mM Tris-HCl, pH 8.0, 10 mM MgCl₂, 50 mM NaCl, 4 mM spermidine, 0.4 μ g/ml BSA, 4 μ l (= 40 U) NheI, 4 μ l (= 40 U) BstEII, in 100 μ l total volume at 37°C.

25 PCR.VIII:

20 mM Tris-Acetate pH 7.5, 20 mM MgAc₂, 100 mM KAc (2x "One-Phor-All" buffer {Pharmacia}), 4 μ l (= 40 U) EcoRI, in 50 μ l total volume at 37°C. After overnight digestion, PCR.VIII-EcoRI was digested with XhoI (overnight at 37°) by

30 the addition of 46 μ l H₂O and 4 μ l (= 40 U) XhoI. PCR.IX:

25 mM Tris-Acetate, pH 7.8, 100 mM KAc, 10 mM MgAc, 1mM DTT (1x "Multi-Core" buffer {Promega}), 4 mM spermidine, 0.4 μ g/ml BSA, 4 μ l (= 40 U) NheI, 4 μ l (= 40 U) BstEII, in 100 μ l total volume at 37°C.

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PCR.X:

50 mM Tris-HCl, pH 8.0, 10 mM MgCl₂, 50 mM NaCl, 4 mM spermidine, 0.4 μ g/ml BSA, 4 μ l (= 40 U) PstI, 4 μ l (= 40 U) EcoRI, in 100 μ l total volume at 37°C.

5

The digested PCR fragments

PCR.I-SacI/BstEII, PCR.II-SfiI/EcoRI,

PCR.III-Nhel/SacI, PCR.IV-XhoI/EcoRI,

PCR.V-Sali/EcoRi, PCR.VI-Sfii/Nhei,

10 PCR.VII-BstEII/NheI, PCR.VIII-XhoI/EcoRI,

PCR.IX-BstEII/NheI, and PCR.X-PstI/EcoRI

were purified on an 1.2% agarose gel using DEAE-membranes NA45 (Schleicher & Schuell) as described by Maniatis et al. The purified fragments were dissolved in $\rm H_2O$ at a

15 concentration of 100-150 ng/ μ l.

Construction of the pGOSA double head expression vectors.

The construction of pGOSA.E (see Figure 8) involved several cloning steps that produced 4 intermediate constructs

- pGOSA.A to pGOSA.D (see Figure 30-34). The final expression vector pGOSA.E and the oligonucleotides in Table 1 above have been designed to enable most specificities to be cloned into the final pGOSA.E construct (Figure 9). The upstream V_H domain can be replaced by any PstI-BstEII V_H
- gene fragment obtained with oligonucleotides PCR.51 and PCR.89 (see Table 1 above). The oligonucleotides DBL.3 and DBL.4 (see Table 1 above) were designed to introduce SfiI and NheI restriction sites in the V_H gene fragments thus allowing cloning of those V_H gene fragments into the SfiI-
- 30 WheI sites as the downstream V_H domain. All V_L gene fragments obtained with oligonucleotides PCR.116 and PCR.90 (see Table 1 above) can be cloned into the position of the V_L .3418 gene fragment as a SacI-XhoI fragment. A complication here however is the presence of an internal
- SacI site in the V_H .3418 gene fragment. Oligonucleotides DBL.8 and DBL.9 (see Table 1 above) are designed to allow cloning of V_L gene fragments into the position of the

V_L.4715 gene fragment as a SalI-NotI fragment. The pGOSA.E derivatives pGOSA.V, pGOSA.S and pGOSA.T with only one or no linker sequences contain some aberrant restriction sites at the new joining points. The V_HA-V_HB construct without a linker lacks the 5' V_HB SfiI site. The V_HB fragment is cloned into these constructs as a BstEII/NheI fragment using oligonucleotides DBL.10 or DBL.11 and DBL.4 (see Table 1 above). The V_LB-V_LA construct without a linker lacks the 5' V_LA SalI site. The V_LA fragment is cloned into these constructs as a XhoI/BcoRI fragment using oligonucleotides DBL.11 and DBL.9 (see Table 1 above),

In the following part of the description the following linkers are mentioned which are also present in the sequence listing:

the (Gly₄Ser)₃ linker, present in SEQ ID NO: 2 as amino acids 109-123 and SEQ ID NO: 10 as amino acids 121-135, the (Gly₄Ser)₃AlaGlySerAla linker (= linkerA), present in SEQ ID NO: 12 as amino acids 121-139, and

20 the (Gly₄Ser)₂Gly₄Val linker (= linkerV), present in SEQ ID NO: 13 as amino acids 108-122.

pGOSA.A

This plasmid is derived from both the Fv.4715-myc construct and the scFv.4715-myc construct.

An SfiI restriction site was introduced between the DNA sequence encoding the $(Gly_4Ser)_3$ linker and the gene fragment encoding the V_L of the scFv.4715-myc construct (see Figure 30). This was achieved by replacing the BstEII-SacI

- fragment of the latter construct by the fragment PCR-I BstEII/SacI (Figure 20) that contains an SfiI site between the DNA encoding the $(Gly_4Ser)_3$ linker and the V_L .4715 gene fragment. The introduction of the SfiI site also introduced 4 additional amino acids (AlaGlySerAla) between the
- (Gly₄Ser)₃ linker and V_L.4715 resulting in a (Gly₄Ser)₃AlaGlySerAla linker (linkerA). The oligonucleotides used to produce PCR-I (DBL.1 and DBL.2,

see Table 1 above) were designed to match the sequence of the framework-3 region of V_H.4715 and to prime at the junction of the DNA encoding the (Gly₄Ser)₃ linker and the V_L.4715 gene fragment, respectively. Thus pGOSA.A can be indicated as: pelB-V_H4715-linkerA-(SfiI)-V_L4715-myc.

pGOSA.B

This plasmid is derived from plasmid Fv.3418 (see Figure 31). The XhoI-EcoRI fragment of plasmid Fv.3418 comprising the 3' end of DNA encoding framework-4 of the V_L including the stop codon was removed and replaced by the fragment PCR-IV XhoI/EcoRI (Figure 23). The oligonucleotides used to produce PCR-IV (DBL.6 and DBL.7, see Table 1 above) were designed to match the sequence at the junction of the V_L and the (Gly₄Ser)₃ linker perfectly (DBL.6), and to be able to prime at the junction of the (Gly₄Ser)₃ linker and the V_H in pUR.4124 (DBL.7). DBL.7 removed the PstI site in the V_H (silent mutation) and introduced a SalI restriction site at the junction of the (Gly₄Ser)₃ linker and the V_H, thereby replacing the last Ser of the linker by a Val residue resulting in a (Gly₄Ser)₂Gly₄Val linker (linkerV). Thus pGOSA.B can be indicated as:

pelB-V_H3418 + pelB-V_L3418-linkerV-(SalI-EcoRI)

25 pGOSA.C

This plasmid contains DNA encoding V_H.4715 linked by the (Gly₄Ser)₃AlaGlySerAla linker to V_H.3418 (see Figure 32), thus: pelB-V_H4715-linkerA-V_H3418.

This construct was obtained by replacing the SfiI-BcoRI fragment from pGOSA.A encoding V_L.4715 by the fragment PCR-II SfiI/EcoRI containing the V_H.3418 gene (see Figure 21). The oligonucleotides used to produce PCR-II (DBL.3 and DBL.4, see Table 1 above) hybridize in the framework-1 and framework-4 region of the gene encoding V_H.3418,

respectively. DBL.3 was designed to remove the PstI restriction site (silent mutation) and to introduce an SfiI restriction site upstream of the V_H gene. DBL.4 destroys

the BstEII restriction site in the framework-4 region and introduces an NheI restriction site downstream of the stopcodon.

5 pGOSA.D

This plasmid contains a dicistronic operon comprising the V_H .3418 gene and DNA encoding V_L .3418 linked by the $(Gly_4Ser)_2Gly_4Val$ linker to V_L .4715 (see Figure 33), thus: $pelB-V_H3418 + pelB-V_L3418-linkerV-V_L4715$.

- This construct was obtained by digesting plasmid pGOSA.B with SalI-EcoRI and inserting the fragment PCR-V SalI-EcoRI (Figure 24) containing the V_L .4715 gene. The oligonucleotides used to obtain PCR-V (DBL.8 and DBL.9, see Table 1 above) were designed to match the nucleotide
- 15 sequence of the framework-1 and framework-4 regions of the $V_L.4715$ gene, respectively. DBL.8 removed the SacI site from the framework-1 region (silent mutation) and introduced a SalI restriction site upstream of the $V_L.4715$ gene. DBL.9 destroyed the XhoI restriction site in the framework-4
- 20 region of the $V_1.4715$ gene (silent mutation) and introduced a NotI and an EcoRI restriction site downstream of the stop codon.

pGOSA.E

- This plasmid contains a dicistronic operon comprising DNA encoding V_H.4715 linked by the (Gly₄Ser)₃AlaGlySerAla linker to V_H.3418 plus DNA encoding V_L.3418 linked by the (Gly₄Ser)₂Gly₄Val linker to V_L.4715 (see Figure 34), thus: pelB-V_H4715-linkerA-V_H3418 + pelB-V_L3418-linkerV-V_L4715.
- Both translational units are preceded by a ribosome binding site and DNA encoding a pelB leader sequence. This plasmid was obtained by a three-point ligation by mixing the vector resulting from pGOSA.D after removal of the V_H3418-encoding PstI-SacI insert with the PstI-NheI pGOSA.C insert
- ontaining V_H.4715 linked to V_H.3418 and the PCR-III

 NheI/SacI fragment (see Figure 22). The remaining PstI-SacI

 pGOSA.D vector contains the 5' end of the framework-1

region of V_H.3418 upto the PstI restriction site and V_L.3418 linked by the (Gly₄Ser)₂Gly₄Val linker to V_L.4715 starting from the SacI restriction site in V_L.3418. The PstI-NheI pGOSA.C insert contains V_H.4715 linked by the (Gly₄Ser)₃-5 AlaGlySerAla linker to V_H.3418, starting from the PstI restriction site in the framework-1 region in V_H.4715. The NheI-SacI PCR-III fragment provides the ribosome binding site and DNA encoding the pelB leader sequence for the V_L.3418-(Gly₄Ser)₂Gly₄Val-V_L.4715 construct. The oligonucleotides DBL.5 and PCR.116 (see Table 1 above) used to generate PCR-III were designed to match the sequence upstream of the ribosome binding site of V_L.4715 in Fv.4715 and to introduce an NheI restriction site (DBL.5), and to match the framework-4 region of V_L.3418 (PCR.116).

15

pGOSA. V

This plasmid is derived from pGOSA.E (see Figure 35) from which the BstEII/NheI fragment containing DNA encoding linkerA-V_H.3418 was excised and replaced by the fragment PCR-VII BstEII/NheI containing the V_H.3418 gene (see Figure 26). The resulting plasmid pGOSA.V contains V_H.3418 linked directly to the framework-4 region of V_H.4715, plus V_L.4715 linked by the (Gly₄Ser)₂Gly₄Val linker to the framework-4 region of V_L.3418, thus:



 $pelB-V_H4715*V_H3418 + pelB-V_L3418-linkerV-V_L4715.$

pGOSA.S

This plasmid is derived from pGOSA.E (see Figure 36) from which the (Gly₄Ser)₂Gly₄Val-V_L4715 *XhoI/Eco*RI fragment was excised and replaced by the fragment PCR-VIII *XhoI/Eco*RI which contains V_L.4715 (see Figure 27). The resulting plasmid pGOSA.S contains V_H.4715 linked by the (Gly₄Ser)₃-AlaGlySerAla linker to V_H.3418 plus V_L.3418 linked directly to the 5' end of the framework-1 region of V_L.4715, thus:

35 $pelb-V_H.4715-linkerA-V_H.3418 + pelb-V_L.3418*V_L.4715.$

pGOSA.T

This plasmid contains a dicistronic operon consisting of $V_H.3418$ directly to the framework-4 region of $V_H.4715$ plus $V_L.3418$ linked directly to the 5' end of the framework-1 region of $V_L.4715$ (see Figure 37). Both transcriptional units are preceded by a ribosome binding site and a pelB leader sequence, thus:

X

pelB-V_H.4715*V_H.3418 + pelB-V_L.3418*V_L.4715.

This construct was obtained by inserting the NheI/EcoRI

fragment of pGOSA.S which contains V_L.3418 linked directly to the 5' end of the framework-1 region of V_L.4715, into the vector pGOSA.V from which the NheI/EcoRI fragment containing V_L.3418 linked by the (Gly₄Ser)₂Gly₄Val linker to V_L.4715 was removed.

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EXAMPLE 6. Detailed description of the preparation of other dicistronic constructs pGOSA.G, and pGOSA.J, pGOSA.Z, pGOSA.AA and pGOSA.AB

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pGOSA.G

This plasmid is an intermediate for the synthesis of pGOSA.J. It is derived from pGOSA.E from which the V_H4715 PstI/BstEII fragment has been excised and replaced by the V_H3418 PstI/BstEII fragment (excised from Fv.3418). The resulting plasmid pGOSA.G (see Figure 38) contains two copies of V_H.3418 linked by the (Gly₄Ser)₃AlaGlySerAla linker, plus V_L.4715 linked by the (Gly₄Ser)₃Gly₄Val linker to the framework-4 region of V_L.3418, thus:

30 $pelB-V_H.3418-linkerA-V_H.3418 + pelB-V_L.3418-linkerV-V_L.4715.$

pGOSA.J

This plasmid contains a dicistronic operon consisting of V_H.3418 linked by the (Gly₄Ser)₃AlaGlySerAla linker to

V_H.4715 plus V_L.3418 linked by the (Gly₄Ser)₇Gly₄Val linker to V_L.4715. Both transcriptional units are preceded by a

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ribosome binding site and a pelB leader sequence (see Figure 39), thus:

 $pelB-V_B.3418-linkerA-V_B.4715 + pelB-V_L.3418-linkerV-V_L.4715$.

5 This construct was obtained by inserting the fragment PCR-VI SfiI/NheI which contains V_H4715 (Figure 25), into the vector pGOSA.G from which the SfiI/NheI V_H3418 fragment was removed.

10 pGOSA.Z

This plasmid is derived from pGOSA.G from which the (Gly₄Ser)₃AlaGlySerAla linker-V_H3418 BstEII/NheI fragment was excised and replaced by the fragment PCR-IX BstEII/NheI which contains V_H.4715 (Figure 28). The resulting plasmid pGOSA.Z (see Figure 40) contains V_H.3418 linked directly to the framework-1 region of V_H.4715, plus V_L.4715 linked by the (Gly₄Ser)₂Gly₄Val linker to the framework-4 region of V_L.3418, thus:

 $pelB-V_H.3418*V_H.4715 + pelB-V_L.3418-linkerV-V_L.4715.$

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pGOSA.AA

This plasmid contains a dicistronic operon consisting of the V_H.3418 linked directly to the 5' end of the framework-1 region of V_H.4715 plus V_L.3418 linked directly to the 5' end of the framework-1 region of V_L.4715. Both transcriptional units are preceded by a ribosome binding site and a pelB leader sequence (see Figure 41). This construct was obtained by inserting the NheI/EcoRI fragment of pGOSA.T which contains V_L.3418 linked directly to the 5' end of the framework-1 region of V_L.4715, into the vector pGOSA.Z from which the NheI/EcoRI fragment containing V_L.3418 linked by the (Gly₄Ser)₂Gly₄Val linker to V_L.4715 was removed, thus:

 $pelB-V_H.3418*V_H.4715 + pelB-V_L.3418*V_L.4715.$

pGOSA.AB

This plasmid is derived from pGOSA.J by a three point ligation reaction (see Figure 42). The SacI/EcoRI insert, containing part of $V_H.3418$ and the full

- 5 (Gly₄Ser)₃AlaGlySerAla linker-V_H.4715 and the V_L.3418-(Gly₄Ser)₂Gly₄Val-V_L.4715 encoding sequences, was removed and replaced by the SacI/SacI pGOSA.J fragment containing the same part of V_H.3418 and the full (Gly₄Ser)₃AlaGlySerAla linker-V_H.4715 and the SacI/EcoRI pGOSA.T fragment
- 10 containing V_L.3418 linked directly to the framework-1 region of V_L.4715 (see Figure 37). The resulting plasmid contains V_H.3418 linked by the (Gly₄Ser)₃AlaGlySerAla linker to the 5' end of the framework-1 region of V_H.4715 plus V_L.3418 linked directly to the 5' end of the framework-1 region of V_L.4715, thus:

 $pelb-V_{H}.3418-linkerA-V_{H}.4715 + pelb-V_{L}.3418*V_{L}.4715$.

EXAMPLE 7. Detailed description of the preparation of monocistronic constructs pGOSA.L and pGOSA.Y, and pGOSA.C, pGOSA.AC and pGOSA.AD

pGOSA.L

This plasmid is derived from pGOSA.E from which the

HindIII/NheI fragment containing DNA encoding V_H.4715.

(Gly₄Ser)₃AlaGlySerAla-V_H.3418 was removed (see Figure 43).

The DNA ends of the vector were made blunt-end using Klenow DNA polymerase and ligated. The resulting plasmid pGOSA.L contains V_L.3418 linked by the (Gly₄Ser)₂Gly₄Val linker to

the 5' end of the framework-1 region of V_L.4715, thus:

pelB-V_L.3418-linkerV-V_L.4715.

pGOSA.Y

This plasmid is derived from pGOSA.T from which the HindIII/NheI fragment containing DNA encoding $V_{\rm H}.4715$ - $V_{\rm H}.3418$ was removed (see Figure 44). The DNA ends of the vector were made blunt-end using Klenow DNA polymerase and

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ligated. The resulting plasmid pGOSA.Y contains $V_L.3418$ linked directly to 5' end of the framework-1 region of $V_L.4715$, thus:

pelB-V_L.3418*V_L.4715.

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The preparation of pGOSA.C was given in Example 5 above; it can be indicated with: pelB-V_H.4715-linkerA-V_B.3418.

pGOSA.X

This plasmid is derived from pGOSA.T from which the NheI/EcoRI fragment containing DNA encoding V_L.3418-V_L.4715 was removed. The DNA ends of the vector were made blunt-end using Klenow DNA polymerase and ligated. The resulting plasmid pGOSA.X (see Figure 45) contains V_H.4715 linked directly to 5' end of the framework-1 region of V_H.3418, thus:

pelB-V_H.4715*V_H.3418.

pGOSA.AC

This plasmid is derived from pGOSA.Z from which the

NheI/EcoRI fragment containing DNA encoding V_L.3418(Gly₄Ser)₂Gly₄Val-V_L.4715 was removed (see Figure 46). The

DNA ends of the vector were made blunt-end using Klenow DNA
polymerase and ligated. The resulting plasmid pGOSA.AC
contains V_H.3418 linked directly to 5' end of the

framework-1 region of $V_{\rm H}$.4715, thus:

 $pelB-V_{H}.3418*V_{H}.4715.$

pGOSA.AD

This plasmid was obtained by inserting the PstI/EcoRI PCR.X

fragment containing DNA encoding V_H.3418-(Gly₄Ser)₃AlaGlySerAla-V_H.4715 (see Figure 29) into the Fv.4715-myc vector
from which the PstI/EcoRI Fv.4715-myc insert was removed
(see Figure 47), thus: pelB-V_H.3418-linkerA-V_H.4715.

35 These monocistronic constructs can be used to transform the same host with two different plasmids or to transform two

different hosts, so that the two $V_{H}{}'s$ in series can be produced separately from the two $V_{L}{}'s$ in series.

Evaluation of the results obtained

- 5 Bifunctional binding activity of GOSA double heads
 In this specification the construction of a two chain
 protein complex is described, in which one of the chains
 consists of two heavy chain V-domains and the other chain
 consists of the two corresponding light chain V-domains.
- The variable domains are linked either directly or through a polypeptide linker. In this specification evidence is provided that these type of molecules ("double heads") contain both antigen binding specificities of the Fv's used to generate these multi-functional antibody fragments.
- 15 Figure 12 shows that GOSA.E can be used to specifically target the enzyme glucose oxidase to several Streptococcus sanguis strains, using antibody fragments derived from hybridomas expressing antibodies directed against these antigens. Figure 12 further shows that the fine specificity of the anti-Streptococcus sanguis scEv 4715 is preserved in
- of the anti-Streptococcus sanguis scFv 4715 is preserved in the GOSA.E double head.

Effect of linkers and relative position of V-domains on double head activity

- After it was shown that the "cross-over double-head" approach (V_HA-V_HB + V_LB-V_LA) yields active bispecific molecules, the importance of the relative position of the V-domains in these constructs was investigated. Both possible positional orientations (GOSA.E = V_HA-LinkerA-V_HB +
- V_LB -LinkerV- V_LA and GOSA.J = V_HB -LinkerA- V_HA + V_LB -LinkerV- V_LA) were constructed and tested for bispecific activity, despite the suggestion obtained by molecular modelling that the binding site formed by the second (downstream/C-terminal) V-domains in the configuration V_HB - V_LA + V_LB - V_LA
- 35 (GOSA.J) was in an unfavourable position for binding to large protein antigens on the surface of cells.
 Surprisingly however, it was found experimentally that the

downstream binding site is in fact accessible. Although the relative position of the heavy chains and the light chains was found to have an effect on the observed reactivity both tested combinations show bispecific activity with the 5 "cross-over" combination (GOSA.E = $V_HA-V_HB + V_LB-V_LA$) exhibiting a higher level of reactivity compared to the combination $V_HB-V_HA+V_LB-V_LA$ (= GOSA.J) as demonstrated for A=anti-Strep and B=anti-Gox.

- 10 Molecular modelling of the $V_HB-V_HA + V_LB-V_LA$ (= GOSA.J) configuration further suggested that, only when the connecting linkers are kept long enough (to span 30 to 35 Å), the protein chains could fold such that both binding sites are fully accessible.
- 15 The "cross-over" configuration: V_HA-V_HB + V_IB-V_IA (GOSA.E) wherein linker length was not critical, was predicted to result in a complex with both binding sites facing in opposite directions, without the restraints suggested for the configuration $V_{\mu}B-V_{\mu}A+V_{\tau}B-V_{\tau}A$ (GOSA.J).
- 20 Removing the flexible polypeptide linker from the V_HA-V_HB chain only had a minimal effect on the ability of the double head in the "cross-over" configuration (GOSA.V = $V_HA*V_HB + V_LB-V_LA$) to bind both S. sanguis and Glucose oxidase. However, removing the flexible polypeptide linker
- 25 from the V_HB-V_HA chain from the molecule in the V_HB-V_HA + V_LB-V_LA configuration (GOSA.Z = $V_HB*V_HA + V_LB-V_LA$) resulted in a dramatic reduction of its ability to bind both S. sanguis and Glucose oxidase.
- In contrast with the double head in the "cross-over" 30 configuration without the flexible polypeptide linker between the two heavy chain domains (GOSA.V), where molecular modelling predicted the resulting molecule to be active, removal of the flexible linker from the V₁B-V₁A chain could not be modelled such that both binding sites
- 35 were fully accessible. ELISA results confirm that the double head in the V_HB-V_HA + V_LB-V_LA configuration without a linker between the two light chain domains (GOSA.AB)

exhibits only minimal S. sanguis and glucose oxidase binding activity. Surprisingly, deletion of the flexible linker from the V_LB-V_LA chain from the double head in the "cross-over" configuration (GOSA.S) only had a small effect 5 on the bispecific activity of the resulting molecule. As expected from the molecular modelling results from the double heads without a flexible linker between the two light chain domains, removal of both the flexible polypeptide linkers from the double head molecules, could 10 not be modelled such that both binding sites were fully accessible. In agreement with the ELISA results obtained with the GOSA.AB construct, the double head in the $V_H B - V_H A$ + V_LB-V_LA configuration without any linkers (GOSA.AA) only exhibits minimal if any S. sanguis and glucose oxidase 15 binding activity. Surprisingly, the double head in the "cross-over" configuration without any linkers (GOSA.T = $V_HA*V_HB + V_LB*V_LA$) still exhibited 25-50% of S. sanguis and glucose oxidase bispecific binding activity when compared to the activity of the double head in the "cross-over" 20 configuration with two linkers (GOSA.E). Thus the conclusion of this work is that modelling can give some indications, but that the computer programmes cannot predict what is possible and what not. Several deviations from the modelling expectations were found. With a 25 paraphrase on an old saying: theories are nice but the experiment is the ultimate proof.

Sensitivity of GOSA double heads

Using an ELISA format it was shown that the sensitivity of the GOSA. E double head is as least as a sensitive as an IgG-glucose oxidase conjugate, as determined by the lowest concentration of *Streptococcus sanguis* antigen immobilised on a solid phase that is still detectable.

GOSA double heads are produced as dimers

FPLC analysis of partially affinity-purified GOSA.E,

GOSA.V, GOSA.S and GOSA.T samples usually gave only one

GOSA double head activity peak as determined by ELISA (Figures 16-19). The position of this peak in the elution pattern indicated that the molecular weight of the GOSA double head is 40-50 kD. Since this molecular weight

5 corresponds to the expected molecular weight of the V_H2 + V_L2 double head dimer, it was concluded that GOSA.E, GOSA.V, GOSA.S and GOSA.T are primarily produced as dimeric molecules. Occasionally an activity peak with an apparent molecular weight of ~200 kD was observed (Figure 16). The

10 presence of glucose oxidase activity in these fractions indicate that these fractions contain GOSA double head complexed with glucose oxidase.

In vitro assembly of GOSA double heads

- 15 It was shown that bifunctionally active dimeric GOSA molecules together in one cell can be produced by translation from one dicistronic messenger (GOSA.E, GOSA.S, GOSA.T, GOSA.V, GOSA.J, GOSA.AB, GOSA.AA and GOSA.Z). In addition high levels of *S. sanguis* and glucose oxidase
- bispecific binding activity is formed when supernatants of cultures producing the separate GOSA subunits are mixed (see Example 7). The effects of linkers and the relative position of the individual $V_{\rm H}$ -domains on the $S.\ sanguis$ and glucose oxidase bispecific binding activity observed in
- 25 these mixing experiments are comparable to the dicistronic constructs.

The constructs described above are summarised in Table 2

30 Table 2A describes intermediate constructs that were not further tested.

Table 2B describes the dicistronic constructs.

Table 2C describes the monocistronic constructs.

- (LiA) stands for the V_H-V_H linker (Gly₄Ser)₃AlaGlySerAla (= linkerA)
- (LiV) stands for the V_L-V_L linker (Gly₄Ser)₂Gly₄Val (= linkerV)

(*) indicates that the two heavy chain domains or the two light chain domains are fused together without a connecting linker.

5

Table 2.

Table 2A

GOSA.A: $V_H.4715-LiA-(SfiI)-V_L.4715-myc$ 10 GOSA.B: $V_H.3418-LiV-V_L.3418-(SalI/EcoRI)$ GOSA.D: $V_H.3418+V_L.3418-LiV-V_L.4715$

GOSA.G: $V_{H}.3418-LiA-V_{H}.3418+V_{L}.3418-LiV-V_{L}.4715$

Table 2B

15 GOSA.E: $V_H.4715-LiA-V_H.3418+V_L.3418-LiV-V_L.4715$ GOSA.S: $V_H.4715-LiA-V_H.3418+V_L.3418*V_L.4715$ GOSA.T: $V_H.4715*V_H.3418+V_L.3418*V_L.4715$

GOSA.V: $V_{H}.4715*V_{H}.3418 + V_{L}.3418-LiV-V_{L}.4715$

20 GOSA.J: $V_{H}.3418-LiA-V_{H}.4715 + V_{L}.3418-LiV-V_{L}.4715$

GOSA.AB: $V_{H}.3418-LiA-V_{H}.4715 + V_{L}.3418*V_{L}.4715$

GOSA.AA: $V_{H}.3418*V_{H}.4715 + V_{L}.3418*V_{L}.4715$

GOSA.Z : $V_{H}.3418*V_{H}.4715 + V_{L}.3418-LiV-V_{L}.4715$

25 Table 2C

 $GOSA.L : V_L.3418-LiV-V_L.4715$

GOSA.Y: $V_L.3418*V_L.4715$

GOSA.AD: $V_H.3418-LiA-V_H.4715$

30 GOSA.AC: V_H.3418*V_H.4715

GOSA.C: V_H.4715-LiA-V_H.3418

GOSA.X : V_H.4715*V_H.3418

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SEQUENCE LISTING

(1) GENERAL INFORMATION:

- (i) APPLICANT:
 - (A) NAME: Unilever PLC
 - (B) STREET: Unilever House, Blackfriars
 - (C) CITY: London
 - (E) COUNTRY: United Kingdom
 - (F) POSTAL CODE (ZIP): EC4P 4BO (GB)
 - (A) NAME: Unilever N.V.
 - (B) STREET: Weena 455
 - (C) CITY: Rotterdam
 - (E) COUNTRY: The Netherlands
 - (F) POSTAL CODE (ZIP): NL-3013 AL
 - (A) NAME: Paul James DAVIS
 - (B) STREET: The Hawthorns, Pavenham Road
 - (C) CITY: Felmersham (Bedfordshire)
 - (E) COUNTRY: United Kingdom
 - (F) POSTAL CODE (ZIP): MK43 7EX (GB)
 - (A) NAME: Cornelis Paul Brik van der LOGT
 - (B) STREET: 1 Bluebell Rise (Peverel Manor Estate)
 - (C) CITY: Rushden (Northamptonshire)
 - (E) COUNTRY: United Kingdom
 - (F) POSTAL CODE (ZIP): NN10 OTU (GB)
 - (A) NAME: Martine Elisa VERHOEIJEN
 - (B) STREET: 1 Tintagel Close (Manor Farm Estate)
 - (C) CITY: Rushden (Northamptonshire)
 - (E) COUNTRY: United Kingdom
 - (F) POSTAL CODE (ZIP): NN10 ONP (GB)
 - (A) NAME: Steve Wilson
 - (B) STREET: 3 Aldenham Close (Goldington)
 - (C) CITY: Bedford,
 - (E) COUNTRY: United Kingdom
 - (F) POSTAL CODE (ZIP): MK41 0FQ (GB)
- (ii) TITLE OF INVENTION: A bifunctional or bivalent antibody fragment analogue
- (iii) NUMBER OF SEQUENCES: 31
- (iv) COMPUTER READABLE FORM:
 - (A) MEDIUM TYPE: Floppy disk
 - (B) COMPUTER: IBM PC compatible
 - (C) OPERATING SYSTEM: PC-DOS/MS-DOS
 - (D) SOFTWARE: PatentIn Release #1.0, Version #1.30 (EPO)
- (v) CURRENT APPLICATION DATA:

APPLICATION NUMBER: EP 95307332.7

- (2) INFORMATION FOR SEQ ID NO: 1:
 - (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 737 base pairs
 - (B) TYPE: nucleic acid
 - (C) STRANDEDNESS: double
 - (D) TOPOLOGY: linear
 - (ii) MOLECULE TYPE: other nucleic acid
 - (A) DESCRIPTION: /desc = "cDNA domains with synthetic linker(s)"

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lis A 30	lsn	Tyr	Leu	Ala	Trp 35	Tyr	Gln	Gln	Lys	Gln 40	Gly	Lys	Ser	Pro	Gln 45	
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Leu I	Leu	Val	Tyr	Tyr	Thr	Thr	Thr	Leu	GCA Ala	Asp	Gly	Val	Pro	Ser	AGG	193
				50					55					60		
									TAT							241
rne s	ser	GIY	Ser 65	GIA	Ser	GIŸ		70	Tyr	Ser	Leu	Lys	11e 75	Asn	Ser	
TIG (	ממי	רושים	GAA	CAT	بلملمك	ccc	እርጥ	ጥስጥ	TAC	TYZT	CAA	CAT	بلعلمك	TCC	እርጥ	289
Leu G	ln	Pro	Glu	Asp	Phe	Gly	Ser	Tyr	Tyr	Сув	Gln	His	Phe	Trp	Ser	209
		80					85					90				
ACT (	CCT	CGG	ACG	TTC	GGT	GGA	GGG	ACC	AAG	CTC	GAG	ATC	AAA	CGG	GGT	337
	95	ALG	Inr	Pne	GrA	100	GIY	THE	Lys	Leu	105	11e	гув	Arg	GIA	
GGA (	GC	GGT	TCA	GGC	GGA	GGT	GGC	אראה	GGC	GGT	GGC	GGA	TCG	CAG	CTC	385
Gly (	Gly	Gly	Ser	Gly	Gly	Gly	Gly	Ser	Gly	Gly	Gly	Gly	Ser	Gln	Val	303
110					115					120					125	
CAG (	CTG	CAG	GAG	TCA	GGA	CCT	GGC	CTG	GTG	GCG	CCC	TCA	CAG	AGC	CTG	433
	Jeu	3111	GIU	130	OTA	PIO	GTA	neu	Val 135	WTS	PTO	ser	GIU	Ser 140	nen	
TCC 1	ATC	AC A	TGC	ልሮሮ	יאנג	ጥፖል	GGG	ጥጥ	TCA	מידיני	ልሮሮ	פפר	ጥልጥ	උදුත	CTV	401
Ser :	Ile	Thr	Cys	Thr	Val	Ser	Gly	Phe	Ser	Leu	Thr	Gly	Tyr	Gly	Val	481
			145					150					155			

DAC	те	Care	י רמר	CAG		רכי א	CCN	220	CCM		C20	maa	~~~	003	ATG	=
Asn	Trp	Val 160	Arg	Gln	Pro	Pro	Gly 165	Lys	Gly	Leu	Glu	Trp 170	Leu	GGA	Met	529
ATT	TGG Trp 175	Gly	GAT Asp	GGA Gly	AAC Asn	ACA Thr 180	GAC Asp	TAT	AAT Asn	TCA Ser	GCT Ala 185	CTC Leu	AAA Lys	TCC Ser	AGA Arg	577
CTG Leu 190	Ser	ATC	AGC Ser	AAG Lys	GAC Asp 195	Asn	TCC Ser	AAG Lys	AGC Ser	CAA Gln 200	GTT Val	TTC Phe	TTA Leu	AAA Lys	ATG Met 205	625
AAC Asn	AGT Ser	CTG Leu	CAC His	ACT Thr 210	GAT Asp	GAC Asp	ACA Thr	GCC Ala	AGG Arg 215	TAC Tyr	TAC Tyr	TGT Cys	GCC Ala	AGA Arg 220	GAG Glu	673
AGA Arg	GAT Asp	TAT Tyr	AGG Arg 225	CTT Leu	GAC Asp	TAC Tyr	TGG Trp	GGC Gly 230	CAA Gln	GGG Gly	ACC Thr	ACG Thr	GTC Val 235	ACC Thr	GTC Val	721
	TCA Ser		TAAC	<b>-</b> CTT												737
(2)	INF	ORMA'	rion	FOR	SEQ	ID 1	10: 2	2 :								
-	(i)	() (1	QUENC A) LE B) TY D) TC	INGTI PE :	i: 24 amir	10 an	ino id		ls							
	1221															
	(11)	) MOI	LECUI	E T	PE:	prot	ein									
			DUENC					BQ I	D NO	): 2:	:					
Asp 1	(xi)	SE		E DE	SCRI	PTIC	)N: S					Ala	Ser _.	Val 15	Gly	
. 1	(xi)	SE(	QUENC	Thr 5	SCR1	PTIC Ser	N: S	Ala	Ser 10	Leu	Ser			15	_	
, 1 Glu	(xi) Ile Thr	SE( Glu Val	QUENC Leu Thr	Thr 5	SCRI Gln Thr	Ser Cys	N: S Pro Arg	Ala Ala 25	Ser 10 Ser	Leu Gly	Ser Asn	Ile	His 30	15 Asn	Tyr	
, l Glu Leu	(xi) Ile Thr	Glu Val Trp 35	Leu Thr 20	Thr 5 Ile Gln	Gln Thr	Ser Cys Lys	Pro Arg Gln 40	Ala Ala 25 Gly	Ser 10 Ser Lys	Leu Gly Ser	Ser Asn Pro	Ile Gln 45	His 30 Leu	15 Asn Leu	Tyr Val	
Glu Leu Tyr	(xi) Ile Thr Ala Tyr 50	Glu Val Trp 35	Leu Thr 20	Thr 5 Ile Gln Thr	Gln Thr Gln	Ser Cys Lys Ala 55	Pro Arg Gln 40 Asp	Ala 25 Gly	Ser 10 Ser Lys Val	Leu Gly Ser Pro	Ser Asn Pro Ser 60	Ile Gln 45 Arg	His 30 Leu Phe	Asn Leu Ser	Tyr Val Gly	
Glu Leu Tyr Ser 65	(xi) Ile Thr Ala Tyr 50 Gly	Glu Val Trp 35 Thr	Leu Thr 20 Tyr	Thr 5 Ile Gln Thr	Gln Thr Gln Leu Gln 70	Ser Cys Lys Ala 55	Pro Arg Gln 40 Asp	Ala 25 Gly Gly Leu	Ser 10 Ser Lys Val	Leu Gly Ser Pro Ile 75	Asn Pro Ser 60 Asn	Ile Gln 45 Arg Ser	His 30 Leu Phe	Asn Leu Ser	Tyr Val Gly Pro 80	
Glu Leu Tyr Ser 65	(xi) Ile Thr Ala Tyr 50 Gly Asp	Glu Val Trp 35 Thr Ser	Leu Thr 20 Tyr Thr Gly	Thr 5 Ile Gln Thr Thr	Gln Thr Gln Leu Gln 70	Cys Lys Ala 55 Tyr	Pro Arg Gln 40 Asp Ser Cys	Ala 25 Gly Gly Leu	Ser 10 Ser Lys Val Lys His 90	Leu Gly Ser Pro Ile 75 Phe	Ser Asn Pro Ser 60 Asn Trp	Ile Gln 45 Arg Ser Ser	His 30 Leu Phe Leu	Asn Leu Ser Gln Pro 95	Tyr Val Gly Pro 80 Arg	
Glu Leu Tyr Ser 65 Glu Thr	(xi) Ile Thr Ala Tyr 50 Gly Asp	Glu Val Trp 35 Thr Ser Phe Gly	Leu Thr 20 Tyr Thr Gly Gly	Thr 5 Ile Gln Thr Thr Ser 85	Gln Thr Gln Leu Gln 70 Tyr	Cys Lys Ala 55 Tyr Tyr Lys	Pro Arg Gln 40 Asp Ser Cys	Ala 25 Gly Gly Leu Gln Glu 105	Ser 10 Ser Lys Val Lys His 90 Ile	Leu Gly Ser Pro Ile 75 Phe	Ser Asn Pro Ser 60 Asn Trp	Ile Gln 45 Arg Ser Ser	His 30 Leu Phe Leu Thr	Asn Leu Ser Gln Pro 95	Tyr Val Gly Pro 80 Arg	
Glu Leu Tyr Ser 65 Glu Thr	(xi) Ile Thr Ala Tyr 50 Gly Asp Phe Gly	Glu Val Trp 35 Thr Ser Phe Gly Gly 115	Leu Thr 20 Tyr Thr Gly Gly 100	Thr 5 Ile Gln Thr Thr Ser 85 Gly	Gln Thr Gln Leu Gln 70 Tyr Thr Ser	Cys Lys Ala 55 Tyr Lys Gly	Pro Arg Gln 40 Asp Ser Cys Leu Gly	Ala 25 Gly Gly Leu Gln Glu 105	Ser 10 Ser Lys Val Lys His 90 Ile	Leu Gly Ser Pro Ile 75 Phe Lys Ser	Ser Asn Pro Ser 60 Asn Trp Arg	Ile Gln 45 Arg Ser Gly Val	His 30 Leu Phe Leu Thr Gly 110 Gln	Asn Leu Ser Gln Pro 95 Gly	Tyr Val Gly Pro 80 Arg Gly	

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Arg Gln Pro Pro Gly Lys Gly Leu Glu Trp Leu Gly Met Ile Trp Gly
165 170 175

Asp Gly Asn Thr Asp Tyr Asn Ser Ala Leu Lys Ser Arg Leu Ser Ile 180 185 190

Ser Lys Asp Asn Ser Lys Ser Gln Val Phe Leu Lys Met Asn Ser Leu 195 200 205

His Thr Asp Asp Thr Ala Arg Tyr Tyr Cys Ala Arg Glu Arg Asp Tyr 210 215 220

Arg Leu Asp Tyr Trp Gly Gln Gly Thr Thr Val Thr Val Ser Ser *
225 230 235 240

### (2) INFORMATION FOR SEQ ID NO: 3:

- (i) SEQUENCE CHARACTERISTICS:
  - (A) LENGTH: 920 base pairs
  - (B) TYPE: nucleic acid
  - (C) STRANDEDNESS: double
  - (D) TOPOLOGY: linear
- (ii) MOLECULE TYPE: other nucleic acid
  - (A) DESCRIPTION: /desc = "cDNA domains with synthetic linker(s)"
- (vii) IMMEDIATE SOURCE:
  - (B) CLONE: HindIII-EcoRI insert Fv.3418
- (ix) FEATURE:
  - (A) NAME/KEY: CDS
  - (B) LOCATION: 36..443
  - (D) OTHER INFORMATION:/product= "pelB-VH3418"
- (ix) FEATURE:
  - (A) NAME/KEY: sig_peptide
  - (B) LOCATION: 36..101
  - (D) OTHER INFORMATION:/product= "pectate lyase"
- (ix) FEATURE:
  - (A) NAME/KEY: mat_peptide
  - (B) LOCATION: 102..440
  - (D) OTHER INFORMATION:/product= "VH3418"
- (ix) FEATURE:
  - (A) NAME/KEY: CDS
  - (B) LOCATION: 495..884
  - (D) OTHER INFORMATION:/product= "pelB-VLA318"
- (ix) FRATURE:
  - (A) NAME/KEY: sig_peptide
  - (B) LOCATION: 495..560
  - (D) OTHER INFORMATION:/product= "pectate lyase"
- (ix) FEATURE:
  - (A) NAME/KEY: mat_peptide
  - (B) LOCATION: 561..881
  - (D) OTHER INFORMATION:/product= "VL3418"
- (xi) SEQUENCE DESCRIPTION: SEQ ID NO: 3:

AAGCTTGCAA ATTCTATTTC AAGGAGACAG TCATA ATG AAA TAC CTA TTG CCT Met Lys Tyr Leu Leu Pro

53

ACC	GCA Ala -15	Ala	GCT Ala	GGA Gly	TTG	Leu -10	Leu	CTC Lev	GCT Ala	GCC Ala	CAA Gln -5	Pro	GCC Ala	ATY Met	GCC Ala	101
CAG Gln 1	val	Gln	CTG Leu	CAG Gln 5	CAG Gln	TCA Ser	GGA Gly	CCI Pro	GAG Glu 10	Leu	GTA Val	AAG Lys	Pro	GGC Gly	GCT Ala	149
TCA Ser	GTG Val	AAG Lys	ATG Met 20	Ser	TGC Cys	AAG Lys	GCT Ala	TCT Ser 25	Gly	TAC	ACA Thr	TTC	ACT Thr	Ser	TAT	197
GII Val	ATG Met	CAC His 35	Trp	GTG Val	AAA Lys	CAG Gln	AAG Lys 40	CCT	GGG Gly	CAG Gln	GGC Gly	CTT Leu 45	GAG Glu	Trp	ATT Ile	245
GGA Gly	TAT Tyr 50	тте	TAT Tyr	CCT Pro	TAC Tyr	AAT Asn 55	gat Asp	GGT Gly	ACT Thr	AAG Lys	TAC Tyr 60	AAT Asn	GAG Glu	AAG Lys	TTC Phe	293
AAA Lys 65	GGC Gly	AAG Lys	GCC Ala	ACA Thr	CTG Leu 70	ACT Thr	TCA Ser	GAC Asp	AAA Lys	TCC Ser 75	TCC Ser	AGC Ser	ACA Thr	GCC Ala	TAC Tyr 80	341
ATG Met	GAG Glu	CTC Leu	AGC Ser	AGC Ser 85	CTG Leu	ACC Thr	TCT Ser	GAG Glu	GAC Asp 90	TCT Ser	GCG Ala	GTC Val	TAT Tyr	TAC Tyr 95	TGT Cys	389
TCA Ser	AGA Arg	CGC Arg	TTT Phe 100	GAC Asp	TAC Tyr	TGG Trp	GGC Gly	CAA Gln 105	GGG Gly	ACC Thr	ACG Thr	GTC Val	ACC Thr 110	GTC Val	TCC Ser	437
TCA Ser	TAA *	TAAC	SAGCT	CAT C	GGAG	CTTG	C AI	rgca/	AATTO	TAT	TTCA	AGG	AGA	CAGT	CAT	493
ser A An Me	" I'G AJ	A TI	AC CI	'A TI	e co	T AC	G GC	A GO	CC GC	TAT T GG	A TT	'G 1"1	TA T	רא כי	rc	<b>493</b> 539
Ser A An Me -2 GCT	TG AA et L 22 GCC	AA TI /B T\ -2 CAA	AC CI FI Le	CA TI	G CC u Pr	T AC	G GC T Al -1	A GO A Al	CC GC la Al	T GG	A TT	G TT tu Le -1	TA TT	TA C.	rc eu	
A AT  Me -2  GCT Ala	TG AM et Ly 22 GCC Ala ATG	CAA Gln -5	AC CT T Le 20 CCA Pro	TA TI	G CC u Pr ATG Met	T AC To Th GCC Ala	GG GC FAI -1 GAC Asp 1	A GO A Al ATC Ile	CC GC la Al GAG Glu ATC	CT GG	ACC Thr 5	G TI tu Le -1 CAG Gln	TA TI	TA C: eu Le CCA Pro	TCT Ser	539
A AT  Me -7  GCT  Ala  TCC  Ser 10	GCC Ala ATG Met	CAA Gln -5 TAT Tyr	AC CTY Let 10 CCA Pro GCA Ala ATT Ile	GCG Ala TCT Ser	ATG Met CTA Leu 15	GCC Ala GGA GGIY	GG GC F Al -1 GAC Asp 1 GAG Glu	AGA ACC	GAG Glu ATC Ile	CTC Leu	ACC Thr 5 ATC	CAG CAG CAG CAG CAG	TA TO	CCA Pro	TCT Ser GCG Ala 25	539 587
A AT  Me -2  GCT Ala  TCC Ser 10  AGT Ser	GCC Ala  ATG Met  CAG Gln  TCT	CAA Gln -5 TAT Tyr GAC Asp	AC CT T Le 20 CCA Pro GCA Ala ATT Ile	GCG Ala TCT Ser AAT ABN 30	ATG Met CTA Leu 15 ACC Thr	GCC Ala GGA Gly TAT Tyr	GG GC TAL GAC Asp 1 GAG Glu TTA Leu	AGA ACC Thr	GAG Glu ATC Ile TGG Trp 35	CTC Leu ACT Thr 20	ACC Thr 5 ATC Ile CAG Gln	CAG Gln ACT Thr	TA TO	CCA Pro AAG Lys CCA Pro 40	TCT Ser GCG Ala 25 GGG Gly	539 587 635
Ser A An Me -2 GCT Ala TCC Ser 10 AGT Ser AAA Lys	GCC Ala  ATG Met  CAG Gln  TCT Ser	CAA Gln -5 TAT Tyr GAC Asp CCC Pro	CCA Pro GCA Ala ATT Ile AAG Lys	GCG Ala TCT Ser AAT Asn 30 ACC Thr	ATG Met  CTA Leu 15 ACC Thr	GCC Ala GGA GIY TAT TYT ATC	GAC ASP 1 GAG GIU TTA Leu TAT Tyr	AGA ACC Thr CGT AGG	GAG Glu ATC Ile TGG Trp 35 GCA Ala	CTC Leu ACT Thr 20 TTC Phe	ACC Thr 5 ATC Ile CAG Gln AGA Arg	CAG Gln ACT Thr CAG Gln TTG Leu	TA TOU LE.O  TCT Ser  TGC Cys  AAA Lys  CTA Leu 55	CCA Pro AAG Lys CCA Pro 40 GAT Asp	TCT Ser GCG Ala 25 GGG Gly	539 587 635
Ser A AT MM -2 GCT Ala TCC Ser 10 AGT Ser AAA Lys STC Val	GCC Ala  ATG Met  CAG Gln  TCT Ser  CCA Pro	CAA Gln -5 TAT Tyr GAC Asp CCC Pro TCA Ser 60 AGC	CCA Pro GCA Ala ATT Ile AAG Lys AGG	GCG Ala TCT Ser AAT Asn 30 ACC Thr	ATG Met  CTA Leu 15 ACC Thr  CTG Leu AGT	GCC Ala GGA GGY TAT Tyr ATC Ile GGC GGY	GG GC  GAC  Asp  GAG  GAG  GAG  TTA  Leu  TAT  Tyr  AGT  65	AGA ACC Thr CGT Arg SGA GGIY	GAG Glu  ATC Ile  TGG Trp 35  GCA Ala  TCT Ser	CTC Leu ACT Thr 20 TTC Phe AAC	ACC Thr 5 ATC Ile CAG Gln AGA Arg	CAG Gln ACT Thr CAG Gln TTG Leu GAT Asp	TA TOU LE.O.  TCT Ser  TGC Cys  AAA Lys  CTA Leu  55  TAT Tyr	CCA Pro AAG Lys CCA Pro 40 GAT Asp TCT Ser	TCT Ser  GCG Ala 25  GGG Gly  GGG GLY  CTC Leu	539 587 635 683

AAA CGG TAA TAATGATCAA ACGGTATAAG GATCCAGCTC GAATTC Lys Arg

920

- (2) INFORMATION FOR SEQ ID NO: 4:
  - (i) SEQUENCE CHARACTERISTICS:
    - (A) LENGTH: 136 amino acids

    - (B) TYPE: amino acid(D) TOPOLOGY: linear
  - (ii) MOLECULE TYPE: protein
  - (xi) SEQUENCE DESCRIPTION: SEQ ID NO: 4:

Met Lys Tyr Leu Leu Pro Thr Ala Ala Ala Gly Leu Leu Leu Leu Ala

Ala Gln Pro Ala Met Ala Gln Val Gln Leu Gln Gln Ser Gly Pro Glu

Leu Val Lys Pro Gly Ala Ser Val Lys Met Ser Cys Lys Ala Ser Gly

Tyr Thr Phe Thr Ser Tyr Val Met His Trp Val Lys Gln Lys Pro Gly

Gln Gly Leu Glu Trp Ile Gly Tyr Ile Tyr Pro Tyr Asn Asp Gly Thr
45 50 55

Lys Tyr Asn Glu Lys Phe Lys Gly Lys Ala Thr Leu Thr Ser Asp Lys

Ser Ser Ser Thr Ala Tyr Met Glu Leu Ser Ser Leu Thr Ser Glu Asp

Ser Ala Val Tyr Tyr Cys Ser Arg Arg Phe Asp Tyr Trp Gly Gln Gly

Thr Thr Val Thr Val Ser Ser * 110

- (2) INFORMATION FOR SEQ ID NO: 5:
  - (i) SEQUENCE CHARACTERISTICS:
    - (A) LENGTH: 130 amino acids
    - (B) TYPE: amino acid
    - (D) TOPOLOGY: linear
  - (ii) MOLECULE TYPE: protein
  - (xi) SEQUENCE DESCRIPTION: SEQ ID NO: 5:

Met Lys Tyr Leu Leu Pro Thr Ala Ala Ala Gly Leu Leu Leu Ala

Ala Gln Pro Ala Met Ala Asp Ile Glu Leu Thr Gln Ser Pro Ser Ser

Met Tyr Ala Ser Leu Gly Glu Arg Ile Thr Ile Thr Cys Lys Ala Ser

Gln Asp Ile Asn Thr Tyr Leu Thr Trp Phe Gln Gln Lys Pro Gly Lys

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Ser Pro Lys Thr Leu Ile Tyr Arg Ala Asn Arg Leu Leu Asp Gly Val

Pro Ser Arg Phe Ser Gly Ser Gly Ser Gly Gln Asp Tyr Ser Leu Thr 60

Ile Ser Ser Leu Asp Tyr Glu Asp Met Gly Ile Tyr Tyr Cys Leu Gln

Tyr Asp Glu Leu Tyr Thr Phe Gly Gly Gly Thr Lys Leu Glu Ile Lys 100

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- (2) INFORMATION FOR SEQ ID NO: 6:
  - (i) SEQUENCE CHARACTERISTICS:
    - (A) LENGTH: 999 base pairs
    - (B) TYPE: nucleic acid
    - (C) STRANDEDNESS: double
    - (D) TOPOLOGY: linear
  - (ii) MOLECULE TYPE: other nucleic acid

(A) DESCRIPTION: /desc = "cDNA domains with synthetic linker(s)"

- (vii) IMMEDIATE SOURCE:
  - (B) CLONE: HindIII-EcoRI insert of Fv.4715-myc
- (ix) FEATURE:
  - (A) NAME/KEY: CDS
  - (B) LOCATION: 40..468
  - (D) OTHER INFORMATION:/product= "pelB-VH4715"
- (ix) FEATURE:
  - (A) NAME/KEY: sig_peptide
  - (B) LOCATION: 40..105
  - (D) OTHER INFORMATION:/product= "pectate lyase"
- (ix) FEATURE:
  - (A) NAME/KEY: mat_peptide
  - (B) LOCATION: 106.465
  - (D) OTHER INFORMATION:/product= "VH4715"
- (ix) FRATURE:
  - (A) NAME/KEY: CDS
  - (B) LOCATION:520..963
  - (D) OTHER INFORMATION:/product= "pelB-VL4715-myc"
- (ix) FEATURE:
  - (A) NAME/KEY: sig_peptide (B) LOCATION:520..585

  - (D) OTHER INFORMATION:/product= "pectate lyase"
- (ix) FRATURE:
  - (A) NAME/KEY: mat_peptide
  - (B) LOCATION: 586. 927
  - (D) OTHER INFORMATION:/product= "VL4715"
- (ix) FRATURE:
  - (A) NAME/KEY: misc RNA
  - (B) LOCATION: 928..960
  - (D) OTHER INFORMATION:/product= "myc-tag"

# (xi) SEQUENCE DESCRIPTION: SEQ ID NO: 6:

(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 6:	
AAGCTTGCAT GCAAATTCTA TTTCAAGGAG ACAGTCATA ATG AAA TAC CTA TTG Met Lys Tyr Leu Leu -22 -20	54
CCT ACG GCA GCC GCT GGA TTG TTA TTA CTC GCT GCC CAA CCA GCG ATG Pro Thr Ala Ala Ala Gly Leu Leu Leu Ala Ala Gln Pro Ala Met -15 -10 -5	102
GCC CAG GTG CAG CTG CAG GAG TCA GGG GGA GAC TTA GTG AAG CCT GGA Ala Gln Val Gln Leu Gln Glu Ser Gly Gly Asp Leu Val Lys Pro Gly 1 5 10 15	150
GGG TCC CTG ACA CTC TCC TGT GCA ACC TCT GGA TTC ACT TTC AGT AGT Gly Ser Leu Thr Leu Ser Cys Ala Thr Ser Gly Phe Thr Phe Ser Ser 20 25 30	198
TAT GCC TTT TCT TGG GTC CGC CAG ACC TCA GAC AAG AGT CTG GAG TGG Tyr Ala Phe Ser Trp Val Arg Gln Thr Ser Asp Lys Ser Leu Glu Trp 35 40 45	246
GTC GCA ACC ATC AGT AGT ACT GAT ACT TAT ACC TAT TAT TCA GAC AAT Val Ala Thr Ile Ser Ser Thr Asp Thr Tyr Thr Tyr Tyr Ser Asp Asn 50 55 60	294
GTG AAG GGG CGC TTC ACC ATC TCC AGA GAC AAT GGC AAG AAC ACC CTG Val Lys Gly Arg Phe Thr Ile Ser Arg Asp Asn Gly Lys Asn Thr Leu 65 70 75	342
TAC CTG CAA ATG AGC AGT CTG AAG TCT GAG GAC ACA GCC GTG TAT TAC Tyr Leu Gln Met Ser Ser Leu Lys Ser Glu Asp Thr Ala Val Tyr Tyr 80 85 90 95	390
TGT GCA AGA CAT GGG TAC TAT GGT AAA GGC TAT TTT GAC TAC TGG GGC Cys Ala Arg His Gly Tyr Tyr Gly Lys Gly Tyr Phe Asp Tyr Trp Gly 100 105 110	438
CAA GGG ACC ACG GTC ACC GTC TCC TCA TAA TAAGAGCTAT GGGAGCTTGC Gln Gly Thr Thr Val Thr Val Ser Ser * 115 120	488
ATGCAAATTC TATTTCAAGG AGACAGTCAT A ATG AAA TAC CTA TTG CCT ACG Met Lys Tyr Leu Leu Pro Thr -22 -20	<b>54</b> 0
GCA GCC GCT GGA TTG TTA TTA CTC GCT GCC CAA CCA GCG ATG GCC GAC Ala Ala Ala Gly Leu Leu Leu Ala Ala Gln Pro Ala Met Ala Asp -15 -5 1	588
ATC GAG CTC ACT CAG TCT CCA TTC TCC CTG ACT GTG ACA GCA GGA GAG Ile Glu Leu Thr Gln Ser Pro Phe Ser Leu Thr Val Thr Ala Gly Glu 5 10 15	636
AAG GTC ACT ATG AAT TGC AAG TCC GGT CAG AGT CTG TTA AAC AGT GTA Lys Val Thr Met Asn Cys Lys Ser Gly Gln Ser Leu Leu Asn Ser Val 20 25 30	684
AAT CAG AGG AAC TAC TTG ACC TGG TAC CAG CAG AAG CCA GGG CAG CCT Asn Gln Arg Asn Tyr Leu Thr Trp Tyr Gln Gln Lys Pro Gly Gln Pro 35 40 45	732
CCT AAA CTG TTG ATC TAC TGG GCA TCC ACT AGG GAA TCT GGA GTC CCT Pro Lys Leu Leu Ile Tyr Trp Ala Ser Thr Arg Glu Ser Gly Val Pro 50 65	780

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924

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GAT Asp	CGC Arg	TTC Phe	ACA Thr	GCC Ala 70	ĀGT Ser	GGA Gly	TCT	GGA Gly	ACA Thr 75	GAT Asp	TTC Phe	ACT Thr	CTC	ACC Thr	ATC Ile
AGC Ser	AGT Ser	GTG Val	CAG Gln 85	GCT Ala	GAA Glu	GAC Asp	CTG Leu	GCA Ala 90	GTT Val	TAT Tyr	TAC Tyr	TGT Cys	CAG Gln 95	Asn	GAT Asp
TAT Tyr	ACT Thr	TAT Tyr 100	CCG Pro	TTC Phe	ACG Thr	TTC Phe	GGA Gly 105	GGG Gly	GGG Gly	ACC Thr	AAG Lys	CTC Leu 110	GAG Glu	ATC Ile	AAA Lys
CGG Arg	GAA Glu 115	CAA Gln	AAA Lys	CTC Leu	ATC Ile	TCA Ser 120	GAA Glu	GAG Glu	GAT Asp	CTG Leu	AAT Asn 125	TAA *	TAA	GATC	AAA
CGG	TAAT	AAG (	SATC	CAGC	rc ga	ATT	2								
	(i)	SEC	)UEN(	E CI	IARAC	TER	STI	CS:							
	(ii)	IOM	A) Li 3) TY D) TC LECUI	INGTI (PE: )POLO	IARAC I: 14 amir XGY: TPE:	3 and according to the second	ino cid car cein	acio		): 7:					
Met -22	(ii) (xi)	(I (I MOI SEC	(UENC	ENGTA (PE: )POLO LE TY LE DE	H: 14 amir XGY: TPE:	3 and according to the second	mino cid car cein	ació	D NO			Leu -10	Leu	Leu	Ala
- 42	(ii) (xi) Lys	MOI SEC	Leu Leu Leu	INGTH (PE: )POLC LE TY LE DE Leu	H: 14 amir XGY: TPB:	3 and according to	nino cid car cein ON: S Ala	ació SEQ I	D NO	Gly	Leu	-10			
-22 Ala	(ii) (xi) Lys Gln -5	MOI SEC Tyr -20	A) LECUI LECUI LECUI LEU Ala	Met	H: 14 amir NGY:  PE: SCRI Pro	and according to the second se	eino ear ein N: S Ala -15	ació SEQ I Ala Gln	D NO Ala Leu	Gly Gln 5	Leu Glu	-10 Ser	Gly	Gly	<b>As</b> p 10
-22 Ala Leu	(ii) (xi) Lys Gln -5 Val	() (E) (E) MOI SEC Tyr -20 Pro	A) LEGUI CECUI CUENC Leu Ala Pro	Met  Gly  178  157  157  157  157  157  157  157	H: 14 amir XGY: XPE: XSCRI Pro Ala	and according protection of the control of the cont	eino ear ein N: S Ala -15 Val	acid SEQ I Ala Gln	D NO Ala Leu Leu 20	Gly Gln 5 Ser	Leu Glu Cys	-10 Ser Ala	Gly Thr	Gly Ser 25	Asp 10 Gly
Ala Leu Phe	(ii) (xi) Lys Gln -5 Val Thr	MOI SEC Tyr -20 Pro Lys	A) LESS TYPE OF THE PROPERTY O	Met Gly Ser	H: 14 amir XGY:  (PE: SSCRI Pro Ala Gly	and according protection of the control of the cont	eino ear ein N: S Ala -15 Val Leu	SEQ I Ala Gln Thr	Ala Leu Leu 20	Gln 5 Ser Val	Leu Glu Cys Arg	-10 Ser Ala Gln	Gly Thr Thr 40	Gly Ser 25 Ser	Asp 10 Gly Asp

Gly Lys Asn Thr Leu Tyr Leu Gln Met Ser Ser Leu Lys Ser Glu Asp
75 80 85

Thr Ala Val Tyr Tyr Cys Ala Arg His Gly Tyr Tyr Gly Lys Gly Tyr 95 100 105

Phe Asp Tyr Trp Gly Gln Gly Thr Thr Val Thr Val Ser Ser *

(2) INFORMATION FOR SEQ ID NO: 8:

- (i) SEQUENCE CHARACTERISTICS:
  - (A) LENGTH: 148 amino acids
    (B) TYPE: amino acid
    (D) TOPOLOGY: linear
- (ii) MOLECULE TYPE: protein

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(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 8:

Met Lys Tyr Leu Leu Pro Thr Ala Ala Ala Gly Leu Leu Leu Ala -22 -20

Ala Gln Pro Ala Met Ala Asp Ile Glu Leu Thr Gln Ser Pro Phe Ser

Leu Thr Val Thr Ala Gly Glu Lys Val Thr Met Asn Cys Lys Ser Gly

Gln Ser Leu Leu Asn Ser Val Asn Gln Arg Asn Tyr Leu Thr Trp Tyr

Gln Gln Lys Pro Gly Gln Pro Pro Lys Leu Leu Ile Tyr Trp Ala Ser

Thr Arg Glu Ser Gly Val Pro Asp Arg Phe Thr Ala Ser Gly Ser Gly

Thr Asp Phe Thr Leu Thr Ile Ser Ser Val Gln Ala Glu Asp Leu Ala

Val Tyr Tyr Cys Gln Asn Asp Tyr Thr Tyr Pro Phe Thr Phe Gly Gly

Gly Thr Lys Leu Glu Ile Lys Arg Glu Gln Lys Leu Ile Ser Glu Glu

Asp Leu Asn 125

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#### (2) INFORMATION FOR SEQ ID NO: 9:

- (i) SEQUENCE CHARACTERISTICS:
  - (A) LENGTH: 924 base pairs
  - (B) TYPE: nucleic acid
  - (C) STRANDEDNESS: double
  - (D) TOPOLOGY: linear
- (ii) MOLECULE TYPE: other nucleic acid

(A) DESCRIPTION: /desc = "cDNA domains with synthetic linker(s)"

- (vii) IMMEDIATE SOURCE:
  - (B) CLONE: HindIII-ECORI insert of scFv.4715-myc
- (ix) FRATURE:
  - (A) NAME/KEY: sig_peptide
  - (B) LOCATION: 40..105
  - (D) OTHER INFORMATION:/product= "pectate lyase"
- (ix) FEATURE:
  - (A) NAME/KEY: mat_peptide
    (B) LOCATION:106..465

  - (D) OTHER INFORMATION:/product= "VH4715"
- (ix) FEATURE:
  - (A) NAME/KEY: misc_RNA
  - (B) LOCATION: 466..510
  - (D) OTHER INFORMATION:/product= "(Gly4Ser)3-linker"
- (ix) FEATURE:
  - (A) NAME/KEY: mat_peptide
  - (B) LOCATION:511..852
  - (D) OTHER INFORMATION:/product= "VL4715"

	(1)	(	A) N B) L D) O	AME/ OCAT	: MOI	853.	.885		oduc	:t= "	myc-	tag"		-		
	(ix	(	ATUR A) N B) L D) O	AME/ OCAT THER	ION:	40 ORMA	888 TION				4715	-myc	n			
	(xi	) SE	QUEN	CE D	ESCR	IPTI	ON:	SEQ	ID N	0: 9	:					
AAG	CTTG	CAT	gcaa	ATTC	та т	TTCA	AGGA	G AC	AGTC			Lys				54
CCT Pro	ACG Thr	GCA Ala -15	Ala	GCT Ala	GGA Gly	TTG Leu	TTA Leu -10	TTA Leu	CTC Leu	GCT Ala	GCC Ala	CAA Gln -5	CCA Pro	GCG Ala	ATG Met	102
GCC Ala	CAG Gln 1	GTG Val	CAG Gln	CTG Leu	CAG Gln 5	GAG Glu	TCA Ser	GGG Gly	GGA Gly	GAC Asp 10	TTA Leu	GTG Val	AAG Lys	CCT Pro	GGA Gly 15	150
GGG Gly	TCC Ser	CTG Leu	ACA Thr	CTC Leu 20	TCC Ser	TGT Cys	GCA Ala	ACC Thr	TCT Ser 25	GGA Gly	TTC Phe	ACT Thr	TTC Phe	AGT Ser 30	AGT Ser	198
TAT Tyr	GCC Ala	TTT Phe	TCT Ser 35	TGG Trp	GTC Val	CGC Arg	CAG Gln	ACC Thr 40	TCA Ser	GAC Asp	AAG Lys	AGT Ser	CTG Leu 45	GAG Glu	TGG Trp	246
GTC Val	GCA Ala	ACC Thr 50	ATC Ile	AGT Ser	AGT Ser	ACT Thr	GAT Asp 55	ACT Thr	TAT Tyr	ACC Thr	TAT Tyr	TAT Tyr 60	TCA Ser	GAC Asp	AAT Asn	294
GTG Val	AAG Lys 65	GGG Gly	CGC Arg	TTC Phe	ACC Thr	ATC Ile 70	TCC Ser	AGA Arg	gac Asp	AAT Asn	GGC Gly 75	AAG Lys	AAC Asn	ACC Thr	CTG Leu	342
TAC Tyr 80	CTG Leu	CAA Gln	ATG Met	AGC Ser	AGT Ser 85	CTG Leu	AAG Lys	TCT Ser	GAG Glu	GAC Asp 90	ACA Thr	GCC Ala	GTG Val	TAT Tyr	TAC Tyr 95	390
TGT Cys	GCA Ala	AGA Arg	CAT His	GGG Gly 100	TAC Tyr	TAT Tyr	GGT Gly	aaa Lys	GGC Gly 105	TAT Tyr	TTT Phe	GAC Asp	TAC Tyr	TGG Trp 110	GGC Gly	438
CAA Gln	GGG Gly	ACC Thr	ACG Thr 115	GTC Val	ACC Thr	GTC Val	TCC Ser	TCA Ser 120	GGT Gly	GGA Gly	GGC Gly	GGT Gly	TCA Ser 125	GGC Gly	GGA Gly	486
GGT Gly	GGC Gly	TCT Ser 130	GGC Gly	GGT Gly	GGC	GGA Gly	TCG Ser 135	GAC Asp	ATC Ile	GAG Glu	CTC Leu	ACT Thr 140	CAG Gln	TCT Ser	CCA Pro	534
TTC Phe	TCC Ser 145	CTG Leu	ACT Thr	GTG Val	ACA Thr	GCA Ala 150	GGA Gly	GAG Glu	AAG Lys	GTC Val	ACT Thr 155	ATG Met	AAT Asn	TGC Cys	AAG Lys	582
TCC Ser 160	GGT Gly	CAG Gln	AGT Ser	CTG Leu	TTA Leu 165	AAC Asn	AGT Ser	GTA Val	AAT Asn	CAG Gln 170	AGG Arg	AAC Asn	TAC Tyr	TTG Leu	ACC Thr 175	630

52	
- mrc TGG	678
TOT ANA CTG TTG ATC TAC	
THE CAG AAG CCA GGG CAG CUT DE LYB Leu Leu 110 190	
TGG TAC CAG CAG AAG CCA GGG CAG CCT CCT AAA CTG TTG ATC TAC TGG TGG TAC CAG CAG AAG CCA GGG CAG CCT CCT AAA CTG TTG ATC TAC TGG TGG TAC CAG CAG AAG CCA GGG CAG CCT CCT AAA CTG TTG ATC TAC TGG 190 185	726
	,
TOT GGA GTC CCT GAT CGC The Thr Ala Ser Gly	
Trp Tyr Gln Gln Lys Flo Grant 185  180  GCA TCC ACT AGG GAA TCT GGA GTC CCT GAT CGC TTC ACA GCC AGT GGA  GCA TCC ACT AGG GAA TCT GGA GTC CCT GAT CGC TTC ACA GCC AGT GGA  Ala Ser Thr Arg Glu Ser Gly Val Pro Asp Arg Phe Thr Ala Ser Gly  200  195  Ala Ser TGT GTG CAG GCT GAA GAC	
	774
Ala Ser Thr Arg Glu Ser 200  Ala Ser Thr Arg Glu Ser 200  TCT GGA ACA GAT TTC ACT CTC ACC ATC AGC AGT GTG CAG GCT GAA GAC  TCT GGA ACA GAT TTC ACT CTC ACC ATC AGC AGT GTG CAG GCT GAA GAC  TCT GGA ACA GAT TTC ACT CTC TTC ACG TTC AC	
TOT GGA ACA GAT TTC ACT CIT THE ILE SET SET VALUE 220	
ser Gly Thr Asp Phe Thr 215	822
210 TAT CCG TTC ACG ITC	
TAT TAC TGT CAG AAT GAT TAY Thr Tyr Pro Pile III	
Ser Gly Thr Asp Phe 1215  210  215  CTG GCA GTT TAT TAC TGT CAG AAT GAT TAT ACT TAT CCG TTC ACG TTC  CTG GCA GTT TAT TAC TGT CAG AAT GAT TAT ACT TAT Tyr Pro Phe Thr Phe  235  CTG GCA GTT TAT TAC TGT CAG AAT GAT TAT ACT TAT CCG TTC ACG TTC  CTG GCA GTT TAT TAC TGT CAG AAT GAT TAT ACT TAT CCG TTC ACG TTC  CTG GCA GTT TAT TAC TGT CAG AAT GAT TAT ACT TAT CCG TTC ACG TTC  CTG GCA GTT TAT TAC TGT CAG AAT GAT TAT ACT TAT CCG TTC ACG TTC  CTG GCA GTT TAT TAC TGT CAG AAT GAT TAT ACT TAT CCG TTC ACG TTC  CTG GCA GTT TAT TAC TGT CAG AAT GAT TAT ACT TAT CCG TTC ACG TTC  CTG GCA GTT TAT TAC TGT CAG AAT GAT TAT ACT TAT CCG TTC ACG TTC  CTG GCA GTT TAT TAC TGT CAG AAT GAT TAT ACT TAT CCG TTC ACG TTC  CTG GCA GTT TAT TAC TGT CAG AAT GAT TAT ACT TAT CCG TTC ACG TTC  CTG GCA GTT TAT TAC TGT CAG AAT GAT TAT ACT TAT CCG TTC ACG TTC  CTG GCA GTT TAT TAC TGT CAG AAT GAT TAT ACT TAT CCG TTC ACG TTC  CTG GCA GTT TAT TAC TGT CAG AAT GAT TAT ACT TAT CCG TTC ACG TTC  CTG GCA GTT TAT TAC TGT CAG AAT GAT TAT ACT TAT CCG TTC ACG TTC  CTG GCA GTT TAT TAC TGT CAG AAT GAT TAT ACT TAT CCG TTC ACG TTC AC	870
	•••
GGA GGG GGG ACC AAG CTC GAG ATC AAA CGG GAA CAA AAA CTC ATC ACC ACC ACC ACC ACC A	
GGA GGG GGG ACC AAG LEU Glu Ile Lys Arg G10 250	604
Gly Gly Thr Ly 245	924
Gly Gly Thr Lys 245 240 245 GAA GAG GAT CTG AAT TAA TAAGATCAAA CGGTAATAAG GATCCAGCTC GAATTC	
CAN GAG GAT CTG AAT TAA TAAGAT	
GAA GAG GAT CLOU ASD * Glu Glu ASD Leu ASD * 260	
250	•
wa. 10:	
(2) INFORMATION FOR SEQ ID NO: 10:	
(2) INFORMATION OF THE STATE OF	
(2) INFORMATION (2013 AMINO ACIDS  (A) LENGTH: 283 AMINO ACIDS  (A) LENGTH: 283 AMINO ACIDS	
(A) mann acid	
(B) TYPE: american (D) TOPOLOGY: linear	
(ii) MOLECULE TYPE: protein (xi) SEQUENCE DESCRIPTION: SEQ ID NO: 10: (xi) SEQUENCE DESCRIPTION: SEQ ID NO: 10:	
(xi) SEQUENCE DESCRIPTION AND Ala Gly Leu	
nro Thr Ala Ala	
Met Lys 192 20 21 Ser Gly Gly Asp	
Ala Gln Val Gln Leu Gln	
Met Lys Tyr Leu Leu Plo -15 -22 -20  Ala Gln Pro Ala Met Ala Gln Val Gln Leu Gln Glu Ser Gly Gly Asp 10  Ala Gln Pro Ala Met Ala Gln Val Gln Leu Gln Glu Ser Gly Gly Asp	
-5 Thr Leu Ser Cys Ala Ini 25	
Ala Gln Pro Ala Met Ala 1  -5  Leu Val Lys Pro Gly Gly Ser Leu Thr Leu Ser Cys Ala Thr Ser Gly  25  Leu Val Lys Pro Gly Gly Ser Leu Thr Leu Ser Cys Ala Thr Ser Asp	
Leu Val Lys Pro Gly 51 20 20 20 20 15  Phe Thr Phe Ser Ser Tyr Ala Phe Ser Trp Val Arg Gln Thr Ser Asp 35 40 35 Thr Asp Thr Tyr Thr	•
Sor Tyr Ala Phe Ser Trp val 40	
Phe Thr Phe Ser Ser 35	
The Ile Ser Ser Thr ASP 1111 35	
Phe Thr Phe Ser Ser 17 35  30  Lys Ser Leu Glu Trp Val Ala Thr Ile Ser Ser Thr Asp Thr Tyr Thr  55  Lys Ser Leu Glu Trp Val Ala Thr Ile Ser Ser Arg Asp Asn	
Lys Ser Leu Glu Trp Val 50 45  Tyr Tyr Ser Asp Asn Val Lys Gly Arg Phe Thr Ile Ser Arg Asp Asn 65  Tyr Tyr Ser Asp Asn Val Lys Gly Arg Phe Thr Ile Ser Arg Asp Asn 65	•
Asp Val Lys Gly Arg Phe 112 70	
Gly Lys Asn Thr Leu Tyr Leu Gln Met Ser Ser Leu Lys Ser Glu Asn	9
60 gin Met Ser Ser Leu Lys Ser 9	0
Cly Lys Asn Thr Leu Tyr Leu 797 185	~
75 Cly Tyr Gly Lys Gly Ty	•
Tyr Cys Ala Arg His Gry 17 100	
Gly Lys Asn Thr Let 180 75 75 Thr Ala Val Tyr Cys Ala Arg His Gly Tyr Tyr Gly Lys Gly Ty 105 Thr Ala Val Tyr Tyr Cys Ala Arg His Gly Tyr Tyr Gly Lys Gly Tyr 100 95 Thr Val Ser Ser Gly Gl	Ly
The Gly Thr Thr Val Thr Val Set 120	
Thr Ala Val Tyr Tyr Gly Gln Gly Thr Thr Val Thr Val Ser Ser Gly Gl  Phe Asp Tyr Trp Gly Gln Gly Thr Thr Val Thr Val Ser Ser Gly Gly  110  Phe Asp Tyr Trp Gly Gln Gly Thr Thr Val Thr Val Ser Ser Gly Gly  110	1u
Phe Asp Tyr Trp Gly Gla 507 115 110 115 Gly Gly Ser Gly	
135	
Gly Gly Ser Gly Gly Gly 130 125  Leu Thr Gln Ser Pro Phe Ser Leu Thr Val Thr Ala Gly Glu Lys 150 140	/al
the Ser Leu Thr Val Thr Ala 317	
Leu Thr Gln Ser Pro Pre 345	
140	

Thr Met Asn Cys Lys Ser Gly Gln Ser Leu Leu Asn Ser Val Asn Gln

Arg Asn Tyr Leu Thr Trp Tyr Gln Gln Lys Pro Gly Gln Pro Pro Lys

Leu Leu Ile Tyr Trp Ala Ser Thr Arg Glu Ser Gly Val Pro Asp Arg

Phe Thr Ala Ser Gly Ser Gly Thr Asp Phe Thr Leu Thr Ile Ser Ser

Val Gln Ala Glu Asp Leu Ala Val Tyr Tyr Cys Gln Asn Asp Tyr Thr

Tyr Pro Phe Thr Phe Gly Gly Gly Thr Lys Leu Glu Ile Lys Arg Glu 235

Gln Lys Leu Ile Ser Glu Glu Asp Leu Asn

- (2) INFORMATION FOR SEQ ID NO: 11:
  - (i) SEQUENCE CHARACTERISTICS:
    - (A) LENGTH: 1706 base pairs
      (B) TYPE: nucleic acid
      (C) STRANDEDNESS: double

  - (ii) MOLECULE TYPE: other nucleic acid

    (A) DESCRIPTION: /desc = "cDNA domains with synthetic linker(s)"
  - (vii) IMMEDIATE SOURCE:
- (B) CLONE: HindIII-ECORI insert of pGOSA.E
  - (ix) FEATURE:
    - (A) NAME/KEY: CDS

    - (D) OTHER INFORMATION:/product= "pelB-VH4715-LiA-VH3418" (B) LOCATION: 40. 864
  - (ix) FEATURE:
    - (A) NAME/KEY: sig_peptide
    - (D) OTHER INFORMATION:/product= "pectate lyase"
  - (ix) FEATURE:
    - (A) NAME/KEY: mat_peptide
    - (B) LOCATION: 106.465
    - (D) OTHER INFORMATION:/product= "VH4715"
  - (ix) FEATURE:
    - (A) NAME/KEY: misc_RNA

    - (B) LOCATION: 466..522
      (D) OTHER INFORMATION:/product= "linkerA (Gly4Ser) 3AlaGlySerAla"
  - (ix) FEATURE:

    - (A) NAME/KEY: mat peptide
      (B) LOCATION:523..861 (D) OTHER INFORMATION:/product= "VH3418"
  - (ix) FEATURE:
    - (A) NAME/KEY: CDS
    - (D) OTHER INFORMATION:/product= "pelB-VL3418-LiV-VL4715" (B) LOCATION:913..1689

	(12,	( <i>)</i>	3) LC	ME/K CATI	ON:9	sig_ 13 RMAT	978		oduct	:= "F	ecta	ite ]	yase	e "	
•	(ix)	(2 (E	3) LC	ME/F	ON:9	mat_ 79	1299	)	oduct	:= "V	/L341	.8 [#]			
	(ix)	(2 (E	3) LC	ME/K CATI HER	ON:1	miso 1300. 1300 1300.	.134 TON:	4 /pro	duct	:= "1	inke	er V			
	(ix)	(2 (E	3) LC	ME/K	[ : NO	mat 1345. DRMAT	.168	16	oduct	:= "V	7 <b>L4</b> 7 1	.5 *			
	(xi)	SEÇ	DENC	E DE	SCRI	PTIC	)N: S	BQ 1	D NC	): 11	. <b>:</b>				•
AAGO	TTGC	AT G	GAA	TTCI	TA TI	TCA	AGGAG	ACA	AGTCA	M		ys I			54
									CTC Leu						102
									GGA Gly						150
									TCT Ser 25						198
									TCA Ser						246
_									TAT Tyr						<b>294</b>
									GAC Asp						342
									GAG Glu						390
	_		_						GGC Gly 105						438
									GGT Gly						486
			Gly						GGT Gly						534

CAA Gln	CAG Gln 145	TCA Ser	GGA Gly	CCT	GAG Glu	CTG Leu 150	GTA Val	AAG Lys	CCT Pro	GGG Gly	GCT Ala 155	Ser	GTG Val	AAG Lys	ATG Met	582
TCC Ser 160	Сув	AAG Lys	GCT Ala	TCT Ser	GGA Gly 165	TAC Tyr	ACA Thr	TTC Phe	ACT Thr	AGC Ser 170	Tyr	GTT Val	ATG Met	CAC His	TGG Trp 175	630
GTG Val	AAA Lys	CAG Gln	AAG Lys	CCT Pro 180	GGG Gly	CAG Gln	GGC Gly	CTT Leu	GAG Glu 185	Trp	ATT Ile	GGA Gly	TAT Tyr	ATT Ile 190	TAT Tyr	678.
			GAT Asp 195												GCC Ala	726
ACA Thr	CTG Leu	ACT Thr 210	TCA Ser	GAC Asp	AAA Lys	TCC Ser	TCC Ser 215	AGC Ser	ACA Thr	GCC Ala	TAC	ATG Met 220	GAG Glu	CTC Leu	AGC Ser	774
AGC Ser	CTG Leu 225	ACC Thr	TCT Ser	GAG Glu	GAC Asp	TCT Ser 230	GCG Ala	GTC Val	TAT Tyr	TAC Tyr	TGT Cys 235	TCA Ser	AGA Arg	CGC Arg	TTT Phe	822
GAC Asp 240	TAC Tyr	TGG Trp	GGC Gly	CAA Gln	GGG Gly 245	ACC Thr	ACC Thr	GTC Val	ACC Thr	GTC Val 250	TCC Ser	TCA Ser	TAA *			864
TAAC	CTAC	GCG (	SAGCT	rgca1	rg CI	\AAT	rcta:	TT	CAAGO	GAGA	CAG	rcat;		Ly:	A TAC S Tyr -20	921
CTA Leu	TTG Leu	CCT Pro	ACG Thr	GCA Ala -15	GCC Ala	GCT Ala	GGA Gly	TTG Leu	TTA Leu -10	TTA Leu	CTC Leu	GCT Ala	GCC Ala	CAA Gln -5	CCA Pro	969
GCG Ala	ATG Met	GCC Ala	GAC Asp 1	ATC Ile	GAG Glu	CTC Leu	ACC Thr 5	CAG Gln	TCT Ser	CCA Pro	TCT Ser	TCC Ser 10	ATG Met	TAT Tyr	GCA Ala	1017
TCT Ser	CTA Leu 15	GGA Gly	GAG Glu	AGA Arg	ATC Ile	ACT Thr 20	ATC Ile	ACT Thr	TGC Cys	AAG Lys	GCG Ala 25	AGT Ser	CAG Gln	GAC Asp	ATT Ile	1065
AAT Asn 30	ACC Thr	TAT Tyr	TTA Leu	ACC Thr	TGG Trp 35	TTC Phe	CAG Gln	CAG Gln	AAA Lys	CCA Pro 40	GGG Gly	aaa Lys	TCT Ser	CCC Pro	AAG Lys 45	- 1113
ACC Thr	CTG Leu	ATC Ile	TAT Tyr	CGT Arg 50	GCA Ala	AAC Asn	AGA Arg	TTG Leu	CTA Leu 55	GAT Asp	GGG Gly	GTC Val	CCA Pro	TCA Ser 60	AGG Arg	1161
TTC Phe	AGT Ser	GGC Gly	AGT Ser 65	GGA Gly	TCT Ser	GGG Gly	CAA Gln	GAT Asp 70	TAT Tyr	TCT Ser	CTC Leu	ACC Thr	ATC Ile 75	AGC Ser	AGC Ser	<b>1209</b>
CTG Leu	GAC Asp	TAT Tyr 80	GAA Glu	gat Asp	ATG Met	GGA Gly	ATT Ile 85	TAT Tyr	TAT Tyr	TGT Cys	CTA Leu	CAA Gln 90	TAT Tyr	GAT Asp	GAG Glu	1257
TTG Leu	TAC Tyr 95	ACG Thr	TTC Phe	GGA Gly	GGG Gly	GGG Gly 100	ACC Thr	AAG Lys	CTC Leu	GAG Glu	ATC Ile 105	AAA Lys	CGG Arg	GGT Gly	GGA Gly	1305
GGC Gly 110	GGT Gly	TCA Ser	GGC Gly	GGA Gly	GGT Gly 115	GGC Gly	TCT Ser	GGC Gly	GGT Gly	GGC Gly 120	GGA Gly	GTC Val	GAC Asp	ATC Ile	GAA Glu 125	1353

56	
- and GTC	1401
GTG ACA GCA GGA GAS VAL	
TOT CCA TIC TCC CTG ACT Thr Ala Gly GIA 140	
CTC ACT CAG 1CT Pro Phe Ser Leu 1112	1449
CTC ACT CAG TCT CCA TTC TCC CTG ACT GTG ACA GCA GGA GAG AAG VAL Leu Thr Gln Ser Pro Phe Ser Leu Thr Val Thr Ala Gly Glu Lys Val 130  ACT ATG AAT TGC AAG TCC GGT CAG AGT CTG TTA AAC AGT GTA AAT CAG  ACT ATG AAT TGC AAG TCC GGT CAG AGT CTG TTA AAC AGT GTA AAT CAG  ACT ATG AAT TGC AAG TCC GGT CAG AGT CTG TTA AAC AGT GTA AAT CAG  TOTAL MET ASIN CYB LYB Ser Gly Gln Ser Leu Leu Asin Ser Val Asin Gln  150	
ACT ATG AAT TGC AAG TCC GGT CAG AGT CTG TTA AAC AGT GTA AAT GAT ATG AAT TGC AAG TCC GGT CAG AGT CTG TTA AAC AGT GTA AAT GAT AAG AGT CTG AAA ASD Ser Val ASD GID 155  Thr Met AsD Cys Lys Ser Gly Gln Ser Leu Leu AsD Ser Val ASD GID 155  Thr Met AsD Cys Lys Ser Gly Gln Ser Leu Leu ASD Ser CCT AAA 145	
ATT TGC AAG TCC GGI Ser Leu Leu ABI 155	
ACT ATO Lys Ser GIF 150	1497
ACT AND AGO LYS LYS SET GIT 150  150  165  150  150  150  150  150	
ACC TIGG TAC CAG CAG AND DETO GLY GIR PRO PLO STA	
AGG AAC TAC TIG ACC TYP TYP GIR GIR BY	1545
AGG AAC TAC TTG ACC TGG TAC CAG CAG AAG CCA GGG CAG CCT CCT AAA  Arg Asn Tyr Leu Thr Trp Tyr Gln Gln Lys Pro Gly Gln Pro Pro Lys  165  165  170  170  170  170  170  170  170  17	1949
THE ACT AGG GAA TOT GGA GIV Val Pro Asp Arg	
ATC TAC TGG GCA TCC ACT Arg Glu Ser Gly Val	
Arg Asn Tyr Leu Thr Trp 165  160  160  CTG TTG ATC TAC TGG GCA TCC ACT AGG GAA TCT GGA GTC CCT GAT CGC  CTG TTG ATC TAC TGG GCA TCC ACT AGG GAA TCT GGA GTC CCT GAT CGC  Leu Leu Ile Tyr Trp Ala Ser Thr Arg Glu Ser Gly Val Pro Asp Arg  180  180  TTC ACT CTC ACC ATC AGC AGT	1593
Leu Leu Ile Tyr Trp Ala 180  180  175  TTC ACA GCC AGT GGA TCT GGA ACA GAT TTC ACT CTC ACC ATC AGC AGT  TTC ACA GCC AGT GGA TCT GGA ACA GAT TTC ACT CTC ACC ATC AGC AGT  TTC ACA GCC AGT GGA TCT GGA ACA GAT TTC ACT CTC ACC ATC AGC AGT  TTC ACA GCC AGT GGA TCT GGA ACA GAT TAT ACT  TTC TCT CAG AAT GAT TAT ACT  TTC TCT CAG AAT GAT TAT ACT	
GCA TCT GGA ACA GAT TIC The Thr Leu Thr Ile Ser 205	
TTC ACA GCC AGT GGA Ser Gly Thr ABP FILE 200	1641
Phe Thr Ala Ser Gly Ser Gly  195  190  GTG CAG GCT GAA GAC CTG GCA GTT TAT TAC TGT CAG AAT GAT TAT ACT  Tyr Thr  GTG CAG GCT GAA GAC CTG GCA GTT TAT TAC TGT Cys Gln Asn Asp Tyr Thr  220  Tal Gln Ala Glu Asp Leu Ala Val Tyr Tyr Cys Gln Asn Asp Tyr Thr  CTG TAA	1041
190 TAT TAC TGT CAG AND ASP TYP THE	
COT GAA GAC CTG GCA GIT TYE TYE CYS GIT AME 220	
GTG CAG GCT GAA GAC CTG GCA GTT TAT TAC TGT CAG AAT GAT TAT TAC  GTG CAG GCT GAA GAC CTG GCA GTT TAT TAC TGT CAG AAT GAT TAT  TYT TYT  220  Val Gln Ala Glu Asp Leu Ala Val Tyr Tyr Cys Gln Asn Asp 220  Val Gln Ala Glu Asp Leu Ala Val Tyr Tyr CTC GAA ATC AAA CGG TAA	1689
VAL GIR AND CTC GAA ATC AAA CGG TAN	
TAT CCG TTC ACG TTC GGA GGG GGG ACC AAG CTC GAA ATC AAA CGG TAA  Tyr Pro Phe Thr Phe Gly Gly Gly Thr Lys Leu Glu Ile Lys Arg  215  215  215  216  217  218  218  219  2219	
TAT CCG TTC ACG TIC Gly Gly Gly Thr hys 233	1706
Tyr Pro Phe Thr 235	1,00
. 225	
TANGCGGCCG CGAATTC	
(2) INFORMATION FOR SEQ ID NO: 12:	
(2) INFORMATION FOR SEE	
(i) SEQUENCE CHARACTERISTICS:  (i) SEQUENCE CHARACTERISTICS:  (ii) SEQUENCE CHARACTERISTICS:	
(i) SEQUENCE CHARACTERISTICS.  (A) LENGTH: 275 amino acids  (A) LENGTH: amino acid	
(A) LENGTH: 2000 acid (B) TYPE: amino acid (B) TYPE: inear	
(D) TOPOLOGI:	
protein	
(ii) MOLECULE TITS. P  (xi) SEQUENCE DESCRIPTION: SEQ ID NO: 12:  (xi) SEQUENCE DESCRIPTION: SEQ ID NO: 12:	
(xi) SEQUENCE DESCRIPTION: SEQ ID NO. 12.  (xi) SEQUENCE DESCRIPTION: SEQ ID NO. 12.  Met Lys Tyr Leu Leu Pro Thr Ala Ala Ala Gly Leu Leu Leu Leu Al  -10  -15  -20  -20  -20  -20  -20  -20  -20  -2	a
(x1) Should have ala Ala Gly Leu Hou -10	_
Tyr Leu Leu Pro 1111 -15	8D
net life -20 con Gln Glu Ser Gly Gly A	10
Met Ala Gln Val Gln Led 5	
Met Lys Tyr Leu Leu Pio Ala -15 -22 -20  Ala Gln Pro Ala Met Ala Gln Val Gln Leu Gln Glu Ser Gly Gly A  Ala Gln Pro Ala Met Ala Gln Val Gln Leu Ser Cys Ala Thr Ser G  Leu Val Lys Pro Gly Gly Ser Leu Thr Leu Ser Cys Ala Thr Ser G  15	ly:
-5 Tell Thr Leu Ser Cys Ala 25	
vol Lys Pro Gly Gly Ser Les 20	n em
Phe Thr Phe Ser Ser Tyr Ala Phe Ser Trp Val Arg Gln Thr Ser 1	HPP
ala Phe Ser Tip 40	
Phe Thr Phe Ser Ser 17 35	Thr
The Ser Ser Thr Asp Inc.	
Tou Glu Trp Val Ala Till 220	
Phe Thr Phe Ser Ser Tyl Add 35  Lys Ser Leu Glu Trp Val Ala Thr Ile Ser Ser Thr Asp Thr Tyr  55  Lys Ser Leu Glu Trp Val Ala Thr Ile Ser Arg Asp	Asn
Val Lys Gly Arg Phe III 70	
Tor Tyr Ser Asp Asn var 65	ASD
Tyr Tyr Ser Asp Ash var 65 60 60 60 61 Leu Gln Met Ser Ser Leu Lys Ser Glu 65 60 60 60 60 60 60 60 60 60 60 60 60 60	90
mor Leu Tyr Leu Gln Met 85	
Gly Lys Ash Line 80	Tyr
75 Arg His Gly Tyr Tyl Gry 2019	5
Gly Lys Asn Thr Leu 191 100 83 75 Thr Ala Val Tyr Tyr Cys Ala Arg His Gly Tyr Tyr Gly Lys Gly 95	
1111 Ann 95	

- Phe Asp Tyr Trp Gly Gln Gly Thr Thr Val Thr Val Ser Ser Gly Gly 110
- Gly Gly Ser Gly Gly Gly Gly Gly Gly Gly Gly Ser Ala Gly Ser 135
- Ala Gln Val Gln Leu Gln Gln Ser Gly Pro Glu Leu Val Lys Pro Gly
- Ala Ser Val Lys Met Ser Cys Lys Ala Ser Gly Tyr Thr Phe Thr Ser
- Tyr Val Met His Trp Val Lys Gln Lys Pro Gly Gln Gly Leu Glu Trp 185
- Ile Gly Tyr Ile Tyr Pro Tyr Asn Asp Gly Thr Lys Tyr Asn Glu Lys
- Phe Lys Gly Lys Ala Thr Leu Thr Ser Asp Lys Ser Ser Ser Thr Ala
- Tyr Met Glu Leu Ser Ser Leu Thr Ser Glu Asp Ser Ala Val Tyr Tyr 220
- Cys Ser Arg Arg Phe Asp Tyr Trp Gly Gln Gly Thr Thr Val Thr Val 250 245

Ser Ser *

- (2) INFORMATION FOR SEQ ID NO: 13:
  - (i) SEQUENCE CHARACTERISTICS:
    - (A) LENGTH: 259 amino acids
    - (B) TYPE: amino acid
    - (D) TOPOLOGY: linear
  - (ii) MOLECULE TYPE: protein
  - (xi) SEQUENCE DESCRIPTION: SEQ ID NO: 13:
- Met Lys Tyr Leu Leu Pro Thr Ala Ala Ala Gly Leu Leu Leu Ala
  -10
  -22 -20
- Ala Gln Pro Ala Met Ala Asp Ile Glu Leu Thr Gln Ser Pro Ser Ser 10
- Met Tyr Ala Ser Leu Gly Glu Arg Ile Thr Ile Thr Cys Lys Ala Ser 25
- Gln Asp Ile Asn Thr Tyr Leu Thr Trp Phe Gln Gln Lys Pro Gly Lys
- Ser Pro Lys Thr Leu Ile Tyr Arg Ala Asn Arg Leu Leu Asp Gly Val
  55
  45
- Pro Ser Arg Phe Ser Gly Ser Gly Ser Gly Gln Asp Tyr Ser Leu Thr
  65
- Ile Ser Ser Leu Asp Tyr Glu Asp Met Gly Ile Tyr Tyr Cys Leu Gln 90
- Tyr Asp Glu Leu Tyr Thr Phe Gly Gly Gly Thr Lys Leu Glu Ile Lys
  105
- Arg Gly Gly Gly Ser Gly Gly Gly Gly Ser Gly Gly Gly Val

PCT/EP96/03605 WO 97/14719

WO 97/14/15	
58	
Asp Ile Glu Leu Thr Gln Ser Pro Phe Ser Leu Thr Val Thr Ala Gly 135 125 126 Ser Leu Leu Asn Ser	
hen He Glu Leu Thr Gln Ser 130	
Asp Ile Glu Leu Thr Gin Son 130  125  Glu Lys Val Thr Met Asn Cys Lys Ser Gly Gln Ser Leu Leu Asn Ser  140  140  130  150  170	
The Live Val The Met Ash Cys Lys See 150	
140 Tyr Gln Gln Lys Pro Gly Gln	
Glu Lys Val Thr Met Ash 145 140  Val Ash Gln Arg Ash Tyr Leu Thr Trp Tyr Gln Gln Lys Pro Gly Gln 160  Val Ash Gln Arg Glu Ser Gly Val	
Val ABN GIL 155 160 160 Thr Arg Glu Ser Gly Val	
155 Low Lew Ile Tyr Trp Ala Ser Inc. 180	
Val Asn Gln Arg Asn 197  160  160  Pro Pro Lys Leu Leu Ile Tyr Trp Ala Ser Thr Arg Glu Ser Gly Val 185  Pro Asp Arg Phe Thr Ala Ser Gly Ser Gly Thr Asp Phe Thr Leu Thr 195  196  197  198  199  199  190  190  190  190  190	
Ser Gly Ser Gly 1 200	
Pro Asp Arg Phe 111 195	
nia Glu Asp Leu Ala var 215	
The Ser Ser Val Gir 2200 210	
The Phe Cly Gly Gly 1111 230	
Asp Tyr Thr Tyr Pro Phe Thr Phe Gly Gly Gly Thr Lys Leu Glu Ile	
220	•
Lys Arg *	
235	
(2) INFORMATION FOR SEQ ID NO: 14:	
(i) SEQUENCE CHARACTERISTICS: (i) SEQUENCE CHARACTERISTICS: (ii) SEQUENCE CHARACTERISTICS:	
(A) IIIII	
(B) TYPE: Bingle	
TYPE: other nucleic action DNA"	
(A) DD2-	
(vii) IMMEDIATE SOURCE:	
(vii) IMMEDIATE SOURCE: (B) CLONE: primer DBL.1 (B) CLONE: primer DBL.1	25
(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 14:	23
CACCATCTCC AGAGACAATG GCAAG	
	•
(2) INFORMATION FOR SEQ ID NO: 15:	
(i) SEQUENCE CHARACTERISTICS:  (i) SEQUENCE CHARACTERISTICS:  (A) LENGTH: 45 base pairs  (A) LENGTH: nucleic acid	
(A) LENGTH: 45 bas of (A) LENGTH: 45 bas of (B) TYPE: nucleic acid (B) TYPE: nucleic acid (B) TYPE: nucleic acid	
(C) STRANDGY: linear	
(D) Torozza acher nucleic acid	
(ii) MOLECULE TYPE: other nucleic acid  (ii) MOLECULE TYPE: other nucleic acid  (A) DESCRIPTION: /desc = "synthetic DNA"	
(vii) IMMEDIATE SOURCE:	
(vii) IMMEDIATE SOURCE: (B) CLONE: primer DBL.2  (B) CLONE: primer DBL.2	41
(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 15:	4
(xi) SEQUENCE DESCRIPTION: DE GAGCGCA GAGCC GAGCGCGAGC TCGGCCGAAC CGGCCGATCC GCCACCGCCA GAGCC	
GAGCGCGAGC TCGGCCGATT	
(2) INFORMATION FOR SEQ ID NO: 16:	
(2) INFORMATION	

(i) SEQUENCE CHARACTERISTICS:
(A) LENGTH: 45 base pairs

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59	
(B) TYPE: nucleic acid (C) STRANDEDNESS: single (D) TOPOLOGY: linear	
(ii) MOLECULE TYPE: other nucleic acid  (A) DESCRIPTION: /desc = "synthetic DNA"	
(vii) IMMEDIATE SOURCE: (B) CLONE: primer DBL.3	
(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 16:	45
(xi) SEQUENCE DESCRIPTION  CAGGATCCGG CCGGTTCGGC CCAGGTCCAG CTGCAACAGT CAGGA	
(2) INFORMATION FOR SEQ ID NO: 17:	
(i) SEQUENCE CHARACTERISTICS.	
(A) IIBNOTTI Jain acid	
(B) TYPE: nucleic single (C) STRANDEDNESS: single (C) STRANDEDNESS: single	
(D) TOPOLOGI.	
(ii) MOLECULE TYPE: other nucleic acid (A) DESCRIPTION: /desc = "synthetic DNA"	
(vii) IMMEDIATE SOURCE:	
(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 17:  CTACATGAAT TCGCTAGCTT ATTATGAGGA GACGGTGACG GTGGTCCCTT GGC	53
CTACATGAAT TCGCTAGCTT ATTATGAGGA GACGG	
(2) INFORMATION FOR SEQ ID NO: 18:	
(2) INFORMATION FOR SECTION (2)	
(i) SEQUENCE CHARACTERISTICS:  (A) LENGTH: 36 base pairs  (A) LENGTH: 316 base pairs	
(A) Individual acid	
(B) TYPE: nucleic single (C) STRANDEDNESS: single (D) TOPOLOGY: linear	
(D) TOPOLOGI.	
(ii) MOLECULE TYPE: other nucleic acid  (A) DESCRIPTION: /desc = "synthetic DNA"	-
(vii) IMMEDIATE SOURCE: (B) CLONE: primer DBL.5	
(B) CHOSEN PROCESSING DESCRIPTION: SEQ ID NO: 18:	36
(XI) SEQUENCE TAGGRAPHIC TATTIC	
(2) INFORMATION FOR SEQ ID NO: 19:	
(i) SEQUENCE CHARACTERISTICS.	
(A) LEROAL ACID	
(C) STRANDSDARD linear	
other nucleic acid	
(ii) MOLECULE TYPE: other nucleic acid  (A) DESCRIPTION: /desc = "synthetic DNA"	
(vii) IMMEDIATE SOURCE: (B) CLONE: primer DBL.6	

WOSHENS	60	
(xi) SEQUENCE DESCRIPT	TION: SEQ ID NO: 19:	23
ACCAAGCTCG AGATCAAACG GGG		•
(2) INFORMATION FOR SEQ I  (i) SEQUENCE CHARACT  (A) LENGTH: 36  (B) TYPE: nucl  (C) STRANDEDNE  (D) TOPOLOGY:	base pairs eic acid SS: single linear	
	GP.	
(vii) IMMEDIATE SOUR (B) CLONE: pr	mer DBL.7 RIPTION: SEQ ID NO: 20:	36
(xi) SEQUENCE DESCR	CCCCACCGCC AGAGCC	
(xi) SECUENCE DE CALLER (XI) S	Coccine	
(2) INFORMATION FOR SE	Q ID NO: 21:	
(i) SEQUENCE CHAP (A) LENGTH: (B) TYPE: no (C) STRANDE	RACTERIS 39 base pairs ucleic acid DNESS: single Y: linear	(
(A) DECU		
(vii) IMMEDIATE S (B) CLONE:	OURCE: primer DBL.8 primer DBL.8	. 39
(xi) SEQUENCE DE	SCRIPTION: SEQ ID NO: 21:	
ATTGGAGTCG ACATCGAAC	CT CACTCAGTCT CCATTCTCC	
(2) INFORMATION FOR	SEQ ID NO: 22:	•
(i) SEQUENCE (A) LENGT (B) TYPE (C) STRAL	HARACISMENT OF THE STATE OF THE	
(M) D=0	TYPE: other nucleic acid CRIPTION: /desc = "synthetic DN	<b>.</b>
(vii) IMMEDIAT (B) CLO	E SOURCE: NE: primer DBL.9	50
(xi) SEQUENCE	DESCRIPTION: SEQ ID NO: 22:	GLCCC
TGAAGTGAAT TCGCGC	GCCGC TTATTACCGT TTGATTTCGA GCTTG	
(2) INFORMATION	FOR SEQ ID NO: 23: TE CHARACTERISTICS: ENGTH: 41 base pairs YPE: nucleic acid	

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61	
(C) STRANDEDNESS: single	
(D) TOPOLOGI:	
(ii) MOLECULE TYPE: other nucleic acid  (ii) MOLECULE TYPE: other nucleic acid  (A) DESCRIPTION: /desc = "synthetic DNA"	
(vii) IMMEDIATE SOURCE:  (B) CLONE: primer DBL.10	
(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 23:	41
CGAATTCGGT CACCGTCTCC TCACAGGTCC ADTT	
(2) INFORMATION FOR SEQ ID NO: 24:	
(i) SEQUENCE CHARACTERISTICS:  (i) SEQUENCE CHARACTERISTICS:  (A) LENGTH: 44 base pairs  (A) LENGTH: pucleic acid	
(A) Librora acid	
(C) STRANDED I inear	
(ii) MOLECULE TYPE: other nucleic acid  (ii) MOLECULE TYPE: other nucleic acid  (iii) MOLECULE TYPE: other nucleic acid  (iv) DESCRIPTION: /desc = "synthetic DNA"	
(A) DECEMENT	
(vii) IMMEDIATE SOURCE: (B) CLONE: primer DBL.11	
CENTERTON: SEQ ID NO.	44
(xi) SEQUENCE DESCRIPTION  COARTTCTCG AGATCAAACG GGACATCGAA CTCACTCAGT CTCC	
(2) INFORMATION FOR SEQ ID NO: 25:	
(i) SEQUENCE CHARACTERISTICS.	
(A) Libridani aria acid	
(C) STRANDBAR linear	
(ii) MOLECULE TYPE: other nucleic acid (iii) MOLECULE TYPE: other nucleic acid (iv) DESCRIPTION: /desc = "synthetic DNA"	
(A) Dabo-	•
(vii) IMMEDIATE SOURCE: (B) CLONE: primer DBL.12	
DESCRIPTION: SEQ ID NO. 22	41
CGAATTCGGT CACCGTCTCC TCACAGGIGC MODE	
(2) INFORMATION FOR SEQ ID NO: 26:	
(i) SEQUENCE CHARACTERISTICS.	
(A) Linda acid	
(C) STRANDADIO 1 linear	
(ii) MOLECULE TYPE: other nucleic acid  (ii) MOLECULE TYPE: other nucleic acid  (A) DESCRIPTION: /desc = "synthetic DNA"	
(A) D3001	

(vii) IMMEDIATE SOURCE:
(B) CLONE: primer PCR.51

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(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 26:	22
AGGTSMAMCT GCAGSAGTCW GG	
(2) INFORMATION FOR SEQ ID NO: 27:  (i) SEQUENCE CHARACTERISTICS:  (A) LENGTH: 32 base pairs	
(C) STRANDER linear (D) TOPOLOGY: linear	
(ii) MOLECULE TYPE: other nucleic acid  (ii) MOLECULE TYPE: other nucleic acid  (A) DESCRIPTION: /desc = "synthetic DNA"	
(vii) IMMEDIATE SOURCE: PCR.89	
(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 27:	32
TGAGGAGACG GTGACCGTGG TCCCTTGGCC	
(2) INFORMATION FOR SEQ ID NO: 28:	
(i) SEQUENCE CHARACTERISTE  (A) LENGTH: 24 base pairs  (B) TYPE: nucleic acid  (C) STRANDEDNESS: single  (D) TOPOLOGY: linear	
(A) Date	
(vii) IMMEDIATE SOURCE: (B) CLONE: primer PCR.90 (B) CLONE: primer PCR.90	24
(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 28:	_
GACATTGAGC TCACCCAGTC TCCA	
(2) INFORMATION FOR SEQ ID NO: 29:	
(i) SEQUENCE CHARACTERISTICS:  (A) LENGTH: 22 base pairs  (B) TYPE: nucleic acid  (C) STRANDEDNESS: single  (D) TOPOLOGY: linear	
(ii) MOLECULE TYPE: other nucleic accumulation (iii) MOLECULE TYPE: other nucl	,
<pre>(vii) IMMEDIATE SOURCE:      (B) CLONE: primer PCR.116  (xi) SEQUENCE DESCRIPTION: SEQ ID NO: 29:</pre>	22
(xi) SEQUENCE DESCRIPTION	
GTTAGATCTC GAGCTTGGTC CC	
(2) INFORMATION FOR SEQ ID NO: 30:  (i) SEQUENCE CHARACTERISTICS:  (i) SEQUENCE CHARACTERISTICS:	
(i) SEQUENCE CHARACTERISTES  (A) LENGTH: 13 amino acids  (B) TYPE: amino acid	

- (D) TOPOLOGY: linear
- (ii) MOLECULE TYPE: protein
- (xi) SEQUENCE DESCRIPTION: SEQ ID NO: 30:

Thr Thr Val Thr Val Ser Ser Gln Val Gln Leu Gln Gln 10

- (2) INFORMATION FOR SEQ ID NO: 31:
  - (i) SEQUENCE CHARACTERISTICS:
    - (A) LENGTH: 12 amino acids
      (B) TYPE: amino acid
      (D) TOPOLOGY: linear
  - (ii) MOLECULE TYPE: protein
  - (xi) SEQUENCE DESCRIPTION: SEQ ID NO: 31:

Lys Leu Glu Ile Lys Arg Asp Ile Glu Leu Thr Gln
10

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BUDAPEST TREATY ON THE INTERNATIONAL RECOGNITION OF THE DEPOSIT OF MICROORGANISMS FOR THE PURPOSES OF PATENT PROCEDURE

INTERNATIONAL FORM

Unilever Research
Colworth Laboratory
Riosciences Division
Colworth House, Sharnbrook
Bedford MK44 ILQ
HAHE AND ADDRESS
OF DEPOSITOR

RECEIPT IN THE CASE OF AN ORIGINAL DEPOSIT ISSUED PURSUANT TO Rule 7.1 by the ISSUED PURSUANT AUTHORITY INTERNATIONAL DEPOSITARY AUTHORITY identified at the bottom of this page

Of Services	
IDENTIFICATION OF THE HIGHOORGANISM	Accession number given by the INTERNATIONAL DEPOSITARY AUTHORITY:
entification reference given by the POSITOR:	NCTC 12916
scherichia coli	
CFy 4715.myc	ONIC DESIGNATION
cFy 4715.myc  1. scientific description and/or proposed taxon	and by:
he microorganism identified under I above was as	Securbous
a scientific description	
a proposed taxonomic designation	
a cross where applicable?	
	identified under I above,
III. RECEIPT AND ACCEPTANCE	s the microorganism letter (date of original deposit)
III. RECEIPT AND ACCEPTANCE  This International Depositary Authority accept which was received by it on 14th October 195	
DEPOSITARY AUTHORITA	plenature(s) of person(s) have Depositary
:: National Collection of Type Cultur	to represent or of authorized
Taboratol	Data: 4/12/95  Bany Hohan  B Holmes Clinical Scientist
Address: Central Public Health Laborator 61 Colindale Avenue	Clinical Scientist
London NW9 SHT	the date on which the status of international deposits of the acquise the acqu
	the date of the fudapest the danget under

here Rule 5.4(d) applies, such date is the date on which the status of international deposition of the status of international deposition authority was acquired; where a deposit made outside the Budapest Treaty atternational depositary authority is converted into a deposit under the of the status of international depositary authority international depositary authority.

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BUDAPEST TREATY ON THE INTERNATIONAL RECOGNITION OF THE DEPOSIT OF MICROGRAMISMS FOR THE PURPOSES OF PATENT PROCEDURE

INTERNATIONAL FORM

Dr P van der Logt lilever Research Sworth Laboratory osciences Division ·Lworth House, Sharnbrook cford MK44 110

VIABILITY STATEMENT rule 10.2 by the issued pursuant to Rule 10.2 by the INTERNATIONAL DEPOSITARY AUTHORITY identified on the following page VIABILITY STATEMENT

WE AND ADDRESS OF THE PARTY WHOM THE VIGORITY STATEMENT IS ISSUED

	II. IDENTIFICATION OF THE MICROORGANISM
Dr P van der/Logt	Accession number given by the INTERNATIONAL DEPOSITANX AUTHORITY:  NCTC 12916:  Date of the deposition of the transfer:
Tame: Ve upone .	lath October: 1995
: VINBILITY STATEMENT	Contified under II above was tested  2. On that date, the said microorganism was
3	
ar loagur viable	or a transfor has been

Indicate the date of the original deposit or, where a new deposit or a transfer has been mide, the most recent relevant date (date of the new deposit or date of the transfer).

The mine cause referred to in Rule 10.2(a) (11) and (111), refer to the most recent viability

1057. Nurk with a gross the applicable box. BP/A/II/12 page 25

CONDITIONS UNDER WHICH THE VINDILITY TEST HAS BEEN PERFORMED.

Nutrient Agar without additions (a)

Agar (17g Bacto-Tryptone, 10g Bacto-Yeast Extract, 5g NaCl, per
11tre) with 100ug/ml ampicillin and 1% glucose (b)

Aerobic incubation, 37°C, 24 hrs

- (a)  $8 \times 10^5$  cfu/ml
- (b) 6 x 10⁵ cfu/ml

Signature(s) of person(s) having the power to represent the international Depositary to represent the authorized official(s): THE EMPATIONAL DEPOSITARY AUTHORITY :.un&: Date: 4/12/95 B Holmes Clinical Scientist #: 0181-000 4400 i.ddiese: Tylen: 8933942 (007434) (1) Par: 0181-203 7874

Fill in if the information has been requested and if the results of the test were negative.

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EUDAPEST TREATY ON THE INTERNATIONAL RECOGNITION OF THE DEPOSIT OF MICROORGANISHS FOR THE PURPOSES OF PATENT PROCEDURE

INTERNATIONAL FORM

TO Dr. P van der Logt
Unilever Research
Colworth Laboratory
Biosciences Division
Colworth House, Sharnbrook
Bedford MK44 11Q
HANE AND ADDRESS
OF DEPOSITOR

RECEIPT IN THE CASE OF AN ORIGINAL DEPOSIT INSUED PURSUANT to Rule 7.1 by the INTERNATIONAL DEPOSITARY AUTHORITY identified at the bottom of this page

entification reference given by the Properties:  Escherichia coli PV 3418	Accossion number given by the INTERNATIONAL DEPOSITARY AUTHORITY: NCIC 12915
DESCRIPTION ALD/OR PROPOSED TAXON	COMPanied by:
i. Scientific description	·
X   a proposed taxonomic designation	
and the second s	s the microorganism identified under I above,
This International Depositary Authority accept which was received by it on 14th October 199  IV. INTERNATIONAL DEPOSITARY AUTHORITY	ALL SAUS
IV. INTERNATIONAL DEFOSION OF Type Culture	to represent the such or state of the state
Address: Central Public Health Laborator 61 Colindale Avenue Lordon NM9 SHT	

Where Rule 6.4(d) applies, such date is the date on which the status of international depositary authority was acquired; where a deposit made outside the Budapest Treaty after the acquisition of the status of international depositary authority is converted into a deposit under the Budapest Treaty, such date is the date on which the microorganism was received by the international depositary authority.

br/n/11/12 page 24

HUDAPEST TREATY ON THE INTERNATIONAL NECOCHITION OF THE DEPOSIT OF MICHOGRAMISMS FOR THE PURPOSES OF PATENT PROCEDURE.

. INTERNATIONAL FORM

. Dr P van der Logt Milever Research Jworth Laboratory iosciences Division olworth House, Sharnbrook edford MK44 ILQ

VIABILITY STATEMENT VIABILITY STATEMENT issued pursuant to Rule 10.2 by the issued pursuant to Rule 10.2 by the INTERNATIONAL DEPOSITARY AUTHORITY identified on the following page

THE AND ADDRESS OF THE PARTY NEED THE VINBILITY STATEMENT IS TISSUED

	II. INENTIFICATION OF THE MICHOGRAMISM
Dr P van der Logt	Accession number given by the INTERNATIONAL PEROSITARY AUTHORITY: NCIC 12915 Date of the deposit or of the transfer: 14th October 1995
	instiffied under II above was tested

The microprogenium identified under II above was tested 2. On that date, the said microorganism was 20th November 1995

Nanka

_ no longer viable

the control with date of the original deposit or, where a new deposit or a transfer has been the control of the criminal deposit or date of the transfer).

In the content to in Rule 10.2(a)(ii) and (iii), refer to the most recent visbility

The specifical across the applicable box.

COMPITIONS UNDER WHICH THE VIABILITY TEST HAS BEEN PERFORMED

Nutrient Agar without additions (a)

Agar (17g Racto-Tryptone, 10g Bacto-Yeast Extract, 5g NaCl, per
litre) with 100ug/sl ampicillin and 17 alucing (b)

Aerobic incubation, 3700, 24 hrs.

- (a) 2 x 10⁶ cfu/nl (b) 1 x 10⁶ cfu/nl

 Natural Collection of Type Collection Central Public Health Laboratory 61 Collection Olst-200 4400 Telephotos 0181-200 4400 Telephotos 9533942 (DEFEND O)	pate: 4/12/95  Balance  Clinical Scientist
 Per: 0181-200 7874	the casults of the test were negative.

Till in if the information has been requested and if the results of the test were negative.

# CLAIMS

- 1. A bispecific or bivalent antibody fragment analogue, which comprises a binding complex containing two polypeptide chains, whereby one polypeptide chain comprises two times a variable domain of a heavy chain  $(V_H)$  in series and the other polypeptide chain comprises two times a variable domain of a light chain  $(V_L)$  in series, and the variable domain of a light chain  $(V_L)$  in series, and the binding complex contains two pairs of variable domains  $(V_H)$  binding complex contains two pairs of variable domains  $(V_H)$  binding complex contains two pairs of variable domains  $(V_H)$  binding complex contains two pairs of variable domains  $(V_H)$  binding complex contains two pairs of variable domains  $(V_H)$  binding complex contains two pairs of variable domains  $(V_H)$  binding complex contains two pairs of variable domains  $(V_H)$  binding complex contains two pairs of variable domains  $(V_H)$  binding complex contains two pairs of variable domains  $(V_H)$  binding complex contains two pairs of variable domains  $(V_H)$  binding complex contains two pairs of variable domains  $(V_H)$  binding complex contains two pairs of variable domains  $(V_H)$  binding complex contains two pairs of variable domains  $(V_H)$  binding complex contains  $(V_H)$
- 2. An antibody fragment analogue according to claim 1, in which one polypeptide chain comprises a first  $V_H$  connected to a second  $V_H$  and the other polypeptide chain comprises a first  $V_L$  connected to a second  $V_L$ .
  - 3. An antibody fragment analogue according to claim 2, in which the two  $V_{H}{}^{\prime}s$  are directly connected to each other without an intermediate peptide linker.
    - 4. An antibody fragment analogue according to claim 2, in which the two  $V_L{}^\prime{}s$  are directly connected to each other without an intermediate peptide linker.
    - 5. An antibody fragment analogue according to claim 3 or claim 4, in which one polypeptide chain comprises a first  $V_H$  directly connected to a second  $V_H$ , and the other polypeptide chain comprises a first  $V_L$  directly connected to a second  $V_L$ .
    - a second  $V_L$ .

      6 An antibody fragment analogue according to claim 2, in which the two  $V_H$ 's are connected to each other by a peptide linker and also the two  $V_L$ 's are connected to each other by a peptide linker appearance appearance appearance are peptide linker comprising at least one amino acid residue.

- Which one polypeptide chain comprises a first  $V_H$  ( $V_{H}$ -A) which one polypeptide chain comprises a first  $V_H$  ( $V_{H}$ -A) and the other polypeptide followed by a second  $V_H$  ( $V_{H}$ -B) and the other polypeptide chain comprises a first  $V_L$  ( $V_L$ -A) followed by a second  $V_L$  chain comprises a first  $V_L$  ( $V_L$ -A) followed by a second  $V_L$  ( $V_L$ -B), and in which the two  $V_H$ 's are connected to each other by a peptide linker ( $Li_H$ ), thus [ $V_H$ -A *  $Li_H$  *  $V_H$ -B], and also the two  $V_L$ 's are connected to each other by a peptide linker ( $Li_L$ ), thus [ $V_L$ -A *  $Li_L$  *  $V_L$ -B], each peptide linker comprising at least 10 amino acid residues.
  - 8. An antibody fragment analogue according to claim 2, in which one polypeptide chain comprises a first  $V_H$  ( $V_{H}$ -A) which one polypeptide chain comprises a first  $V_H$  ( $V_{H}$ -A) with or without a connecting followed by a second  $V_H$  ( $V_{H}$ -B) with or without a connecting peptide linker ( $\text{Li}_H$ ), thus [ $V_{H}$ -A * ( $\text{Li}_H$ ) *  $V_{H}$ -B], and the other polypeptide chain comprises a first  $V_L$  ( $V_{L}$ -A) preceded by a second  $V_L$  ( $V_{L}$ -B) with or without a connecting peptide linker ( $\text{Li}_L$ ), thus [ $V_{L}$ -B * ( $\text{Li}_L$ ) *  $V_{L}$ -A].
    - 9. An antibody fragment analogue according to claim 1, in which the two variable domains are different resulting in a bispecific antibody fragment analogue.
      - 10. An antibody fragment analogue according to claim 1, in which the specificities A and B are the same resulting in a bivalent antibody fragment analogue.
      - 11. Use of an antibody fragment analogue according to claim 1, in immunoassays including diagnostic techniques, in agglutination assays, in a purification method, for compositions suitable for therapy, or in other methods in which immunoglobulins or fragments thereof can be used.
        - 12. A process for producing an antibody fragment analogue according to any one of claims 1-10, which comprises
        - (1) transforming a host by incorporating into that host a DNA encoding the two  $V_{\text{H}}{}^{\prime}s$  in series with or without a

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connecting peptide linker and a DNA encoding the two  $V_{L}{}^{\prime}s$  in series with or without a connecting peptide linker,

- (2) culturing such transformed host under conditions whereby the connected  $V_{H}{}'s$  and the connected  $V_{L}{}'s$  are
- (3) allowing the two connected  $V_{H}{}'s$  and the two connected  $V_{L}{}^{\prime}s$  to combine to each other under formation of a formed, and double head antibody fragment analogue, and
- (4) optionally collecting the double head antibody fragment analogue.
- A process for producing an antibody fragment analogue according to any one of claims 1-10, which
- (1) transforming a first host by incorporating into that first host a DNA encoding the two  $V_{\mbox{\scriptsize H}}{}'s$  in series with or comprises without a connecting peptide linker,
  - (2) transforming a second host by incorporating into that second host a DNA encoding the two  $V_{\text{L}}{}^{\prime}\text{s}$  in series with or without a connecting peptide linker,
  - (3) culturing the first and second transformed host under conditions whereby the connected  $V_{\mbox{\scriptsize H}}{}'s$  and the connected  $V_{l}$ 's, respectively, are formed,
  - (4) optionally collecting separately the two connected  $\boldsymbol{V}_{H}{'}\boldsymbol{s}$  and the two connected  $\boldsymbol{V}_{L}{'}\boldsymbol{s},$  and
  - (5) combining the connected  $V_{H}{}'s$  and the connected  $V_{L}{}'s$ under conditions that they can form a double head antibody fragment analogue.
    - A process according to claim 12 or 13, in which the host is selected from the group consisting of prokaryotic micro-organisms comprising Gram-negative bacteria (e.g. E. coli) and Gram-positive bacteria (e.g. B. subtilis or lactic acid bacteria), lower eukaryotic microorganisms comprising yeasts (e.g. belonging to the genera Saccharomyces, Kluyveromyces, Pichia, and Hansenula) and moulds (e.g. belonging to the genera Aspergillus,

Neurospora or Trichoderma), and higher eukaryotic organisms (e.g. plants) or cell cultures thereof (e.g. hybridoma's).

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Fig.1.

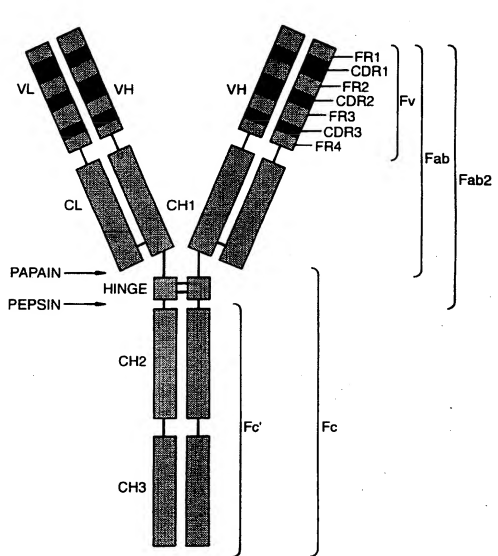
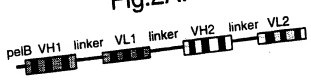


Fig.2A.



# Fig.2B.

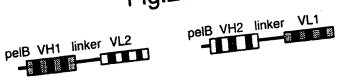
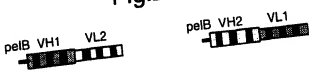
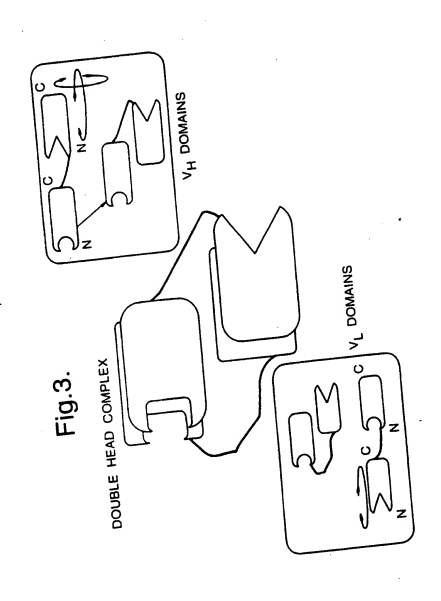


Fig.2C.





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	e E
S L S A S LOPETICTGG	GAGA
Fig.4.  B L T O S P A S L S A S V  GAATTCGGCCGACATCGAGCTCACCCAGTCTCCAGCCTCCCTTTCTGCGTCTGTGG  Saci  FCORI  SACI  N. L H. N. Y L A N.	6.0
CAATTCGGCCGACATC GAGCI	A Ö
ECORI G N I H N I TATITAGCATGG	TATCA
C R A STOCCAATATICACAATA	120
T V T L A AACTGTCACCATCACATGTCGAGCAAGTGGGGAAGTGGGGAAGTGGGGAAGTGGGGAAGTGGGGAAGTGGGGGAAGTGGGGGAAGTGGGGGAAGTGGGGGG	GATGG
AACTGTCACCATCATCATCATCATCATCATCATCATCATCATCAT	180
Q K Q GAAAATCTCCTCAGCTCCTGG	N S
GCAGAACAGGGAAAATCTCCTCAGCICCOGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGG	AACAG
G G S G S G S G S G S G S G S G S G S G	240
V P S R T P	R I
GCAGAACAGGGAAL  V P S R F S G S G T Q Y S L K T  V P S R F S G S G T Q Y S L K T  TGTGCCATCAAGGTTCAGTGGCAGTGGATCAGGAACACATATTCTCTCAAGATC  TGTGCCATCAAGGTTCAGTGGCAGTGGATCAGGAACACATTTTTGGAGTACTCC  CCTGCAACCTGAAGATTTTTGGGAGTTATTACTGTCAACATTTTTTGGAGTACTCC  CCTGCAACCTGAAGATTTTTGGGAGTTATTACTGTCAACATTTTTTTT	300
L Q P B D TITTGGGAGTTATTACIGICA	GG
CCTGCAACCTGAAGAT	CCTGG
F G G G T K L B I K R G G G G G G G G G G G G G G G G G G	360
F G G G T AGCTCGAGATCAAACGGGG	
GTTCGGTGGAGGGACCAAGCTCGAGATCAGACTCGAGATCAGGACCTGGCCTGGCCTGGCCTGGCCTGCAGGAGTCAGGACCTGGCCTGGCCTGGCCTGCAGGAGTCAGGACCTGGCCTGGCCTGGCAGGAGTCAGGACCTGGCCTGGCCTGGCCTGGCAGGAGTCAGGACCTGGCCTGGCAGGAGTCAGGACCTGGCCTGGCAGGAGTCAGGACCTGGCCTGGCCTGGCAGGAGTCAGGACCTGGCCTGGCCTGGCAGGACCTGGCCTGGCAGGAGTCAGGACCTGGCCTGGCCTGGCCTGGCCTGGCCTGGCAGGAGTCAGGACCTGGCCTGCAGGAGCTGCAGGAGCTGGCCTGGCCTGGCCTGGCCTGCAGGAGCTGCAGGAGCTGGCCTGGCCTGGCCTGGCCTGGCCTGGCCTGGCCTGGCCTGGCCTGGCCTGGCCTGGCCTGGCCTGGCCTGGCCTGGCCTGGCCTGGCCTGGCCTGGCCTGGCCTGGCCTGGCCTGGCCTGGCCTGGCCTGGCCTGGCCTGGCCTGGCCTGGCCTGGCCTGGCCTGGCCTGGCCTGGCCTGGCCTGGCCTGGCCTGGCCTGCCTGCAGGAGCCTGGCCTGGCCTGGCCTGGCCTGGCCTGGCCTGGCCTGGCCTGGCCTGGCCTGCCCTGCCTGCCTGCCTGCCTGCCTGCCTGCCTGCCTGCCCTGCCTGCCCTGCCCTGCCCTGCCCTGCCCTGCCCTGCCCTGCCCTGCCCTGCCCTGCCCTGCCCTGCCCTGCCCTGCCCTGCCCTGCCCTGCCCTGCCCTGCCCTGCCCTGCCCTGCCCTGCCCTGCCCTGCCCTGCCCTGCCCTGCCCTGCCCTGCCCTGCCCTGCCCTGCCCTGCCCTGCCCTGCCCTGCCCTGCCCTGCCCTGCCCTGCCCTGCCCTGCCCTGCCCTGCCCTGCCCTGCCCTGCCCTGCCCTGCCCTGCCCTGCCCTGCCCTGCCCTGCCCTGCCCTGCCCTGCCCTGCCCTGCCCTGCCCTGCCCTGCCCTGCCCTGCCCTGCCCTGCCCCTGCCCTGCCCTGCCCTGCCCTGCCCTGCCCTGCCCTGCCCTGCCCTGCCCTGCCCTGCCCTGCCCTGCCCTGCCCTGCCCTGCCCTGCCCTGCCCCTGCCCTGCCCCTGCCCTGCCCCTGCCCCTGCCCCCC	V A F
S G G G G STATEGEAGGTGCAGCTGCAGGAGAGAGAGAGAGAGAGAGAGAGAGA	Y G V
S G G G S Q V Q L Q B S G P G L  S G G G S Q V Q L Q B S G P G L  CTCTGGCGGTGCGGATCGCAGGTGCAGGTGCAGGAGTCAGGACCTGGCCTGG  Pst1  S Q S L S I T C T V S G F S L T G  CTCAGAGAGCCTGTCCATCACATGCACCGTCTCAGGGTTCTCATTAACCGGCT	ratggtg1
TAS TARGET CACCATC TO CACATC TO CACCATC TO CACCATC TO CACCATC TO CACCATC TO CACCATCATC TO CACCATC TO CACCATC TO CACCATC TO CACCATC TO CACCATC TO CACCATC T	
CTCTGGCGTTGCCGATCGCAGGTTGCAGGTTCTCAGTTAACCGGC  S Q S L S I T C T V S G F S L T G  CTCACAGAGCCTGTCCATCACATGCACCGTCTCAGGGTTCTCATTAACCGGC  CTCACAGAGCCTGTCCATCACATGCACCGTCTCAGGGTTCTCATTAACCGGC  N W V R Q P P G K G L B W L G M I	W G D
G L B W L GGAATGATT	TGGGGTGA
CTCACAGAGCCTO2  CTCACAGAGCCTO2  N W V R Q P P G K G L B W L G M I  N W V R Q P P R G K G L B W L G M I  N W V R Q P P R G K G L B W L G M I  N W V R Q P R G M I I I I I I I I I I I I I I I I I I	. 540
K S R L S TOATCAGCAA	GGACAACTC
N W K AAACTGGGTTCGCCAGCCTCCAGGAAAGGGTOO AAACTGGGTTCGCCAGCCTCCAGGAAAGGGTOO AAACTGGGTTCGCCAGCCAGCCAAATCCAGACTGAGCATCAGCAAATCCAGACTGAGCATCAGCAAATCCAGACTGAGCATCAGCAAATCCAGACTGAGCATCAGCAAATCCAGACTGAGCATCAGCAAATCCAGACTGAGCATCAGCAAATCCAGACTGAGCATCAGCAAATCCAGACTGAGCATCAGCAAATCCAGACTGAGCATCAGCAAATCCAGACTGAGCATCAGCAAATCCAGACTGAGCATCAGCAAATCCAGACTGAGCATCAGCAAATCCAGACTGAGCATCAGCAAATCCAGACTGAGCATCAGCAAATCCAGACTGAGCATCAGCAAATCCAGACTGAGCATCAGCAAATCCAGACTGAGCAAAATCCAGACTGAGCAAAATCCAGACTGAGCAAAATCCAGACTGAGCAAAATCCAGACTGAGCAAAATCCAGACTGAGCAAAATCCAGACTGAGCAAAATCCAGACTGAGCAAAATCCAGACTGAGCAAAATCCAGACTGAGCAAAATCCAGACTGAGCAAAATCCAGACTGAGCAAAAATCCAGACTGAGCAAAAATCCAGACTGAGCAAAAAAAA	800
G N T TGGAAACACAGACTATAATTCAGCTCTCAGG TGGAAACACAGACTATAATTCAGCTCTCAGGCACACTGATGACACAGGC  K S Q V F L K M N S L H T D D T A  K S Q V F L K M N S L H T D D T A  CNACAGCCAAGTTTTCTTAAAAATGAACAGTCTGCACACTGATGACACAGC	R Y Y
TGGAAACACAGACTTAAAAAATGAACAGTCTGCACACTGATGACACAGCCCAGCCCAGGCCAAGAGCCAAGTTTTCTTAAAAAATGAACAGTCTGCACACTGATGACAGCCAAGAGCCAAGTTTTCTTAAAAAATGAACAGTCTGCACACTGATGACAGCCAAGAGCCAAGTTTTCTTAAAAAATGAACAGTCTGCACACTGATGACACAGCCAAGAGCCAAGTTTTCTTAAAAAATGAACAGTCTGCACACTGATGACACAGCCAAGAGCCAAGTTTTCTTAAAAAATGAACAGTCTGCACACTGATGACAGCCAAGAGCCAAGTTTTCTTAAAAAATGAACAGTCTGCACACTGATGACAGCCAAGAGCCAAGTTTTCTTAAAAAAATGAACAGTCTGCACACTGATGACACAGCCAAGTTTTTCTTAAAAAAATGAACAGTCTTGCACACTGATGACACAGCCAGC	CAGGTACTA
ANTGAACAGTCTGCACACTGATGATCA	
CARGAGCCAAGTTTTCTTAAAAATGAACCACCACGAGCCAAGGGACCACCACCAAGGGACCAAGGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAG	TVTV
T D Y W G O G	CGGTCACCGT
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CAAGAGCCAAGTTT  CAARERDYRLDYWGQGGCCAAGGGACCA  CTGTGCCAGAGAGAGAGAGATTATAGGCTTGACTACTGGGGCCAAGGGACCA	
contact the second seco	
DIELTOSPAS GGGTKLEIKR ULLys Linker VHLys	
DIELTOSPAS GGGGGGGGGGGGG GGGGGGGGGGGGGGGGGGGGG	
QVOLDESGEG GOGLET VICE SHEET (RULE 26)	
THE SHEET (NOLL 24)	

M K Y L L P T A AAGCTTGCAAATTCTATTTCAAGGAGACAGTCATAATGAAATACCTATTGCCTACGGCAG Fig.5. AAGLLLAAQPAMAQVQLQQ CCGCTGGATTGTTATTACTCGCTGCCCAACCAGCGATGGCCCAGGTGCAGCTGCAGCAGT pelB S G P B L V K P G A S V K M S C X A S G LEADER CAGGACCTGAGCTGGAAAGCCTGGGGCTTCAGTGAAGATGTCCTGCAAGGCTTCTGGAT Y T F T S Y V M H W V K O K P G O G L K ACACATTCACTAGCTATGTTATGCACTGGGTGAAACAGAAGCCTGGGCAGGGCCTTGAGT WIGNIVPYNDGTKYNEKFRG GGATTGGATATTTTATCCTTACAATGATGGTACTAAGTACAATGAGAAGTTCAAAGGCA KATLTSDKSSTAYHELS.S.L. VH3418 AGGCCACACTGACTTCAGACAAATCCTCCAGCACAGCCTACATGGAGCTCAGCAGCCTGA TSBDSAVYYCSRRPDYWGQO CCTCTGAGGACTCTGCGGTCTATTACTGTTCAAGACGCTTTGACTACTGGGGCCAAGGGA TTVTVSS MKYLLPTAAAGLLLL AGGAGACAGTCATAATGAAATACCTATTGCCTACGGCAGCCGCTGGATTGTTATTACTCG LEADER A A Q P A M A D I B L T Q S P S S M Y A CTGCCCAACCAGCGATGGCCGACATCGAGCTCACCCAGTCTCCATCTTCCATGTATGCAT LGERITICKASQDINTYL CTCTAGGAGAGAATCACTATCACTTGCAAGGCGAGTCAGGACATTAATACCTATTTAA WFQQKFGKSPKTLIYRANE CCTGGTTCCAGCAGAAACCAGGGAAATCTCCCAAGACCCTGATCTATCGTGCAAACAGAT VL3418 L. L. D. G. V. P. S. R. F. S. G. S. G. S. G. Q. D. Y. S. L. TGCTAGATGGGGTCCCATCAAGGTTCAGTGGCAGTCGGGCAAGATTATTCTCTCA TISSLDYBDMGIYYCLBQYDE CCATCAGCAGCCTGGACTATGAAGATATGGGAATTTATTATTGTCTACAATATGATGAGT LYTFGGGTKLEIKR TGTACACGTTCGGAGGGGGGGCCAAGCTCGAGATCAAACGGTAATAATGATCAAACGGT ATAAGGATCCAGCTCGAATTC

F	ig.6.
pelB LEADER	AAGCTTGCATGCAAATTCTATTTCAAGGAGACAGTCATAATGAAATACCTATTGCCTACG
	A A A G L L L A A Q P A M A Q W C L O GCAGCCGCTGGATTGTTATTACTCGCTGCCCAACCAGCGATGGCCCAGGTGCAGCTGCAG
VH4715	E 6 G G D L V K P K G S L T L S C A T S GAGTCAGGGGGAGACTTAGTGAAGCCTGGAGGGTCCCTGACACTCTCTGTGCAACCTCT
	G F T F G S T A F S W V R O T S D K S T GGATTCACTTCAGTAGTTATGCCTTTTCTTGGGTCCGCCAGACCTCAGACAAGAGTCTG
	E W V A T I S E T D T Y T Y Y S D N V K GAGTGGGTCGCAACCATCAGTAGTACTGATACTTATACCTATTATTCAGACAATGTGAAG
	G R F T I S R D N G K N T L Y L Q M S S GGGCGCTTCACCATCTCCAGAGACAATGGCAAGAACACCCTGTACCTGCAAATGAGCAGT
	L K S B D T A V Y Y C A R H G Y Y G K G CTGAAGTCTGAGGACACAGCCGTGTATTACTGTGCAAGACATGGGTACTATGGTAAAGGC
pelB LEADER	Y F D Y N G O G T T V T V S S  TATTTTGACTACTGGGGCCAAGGGACCACGGTCACCGTCTCCTCATAATAAGAGCTATGG
	M K Y L L P T GAGCTTGCATGCAAATTCTATTTCAAGGAGACAGTCATAATGAAATACCTATTGCCTACG
	A A A G L L L A A Q P A M A D I E L T GCAGCCGCTGGATTGTTATTACTCGCTGCCCAACCAGCGATGGCCGACATCGAGCTCACT
	Q S P F S L T V T A G E K V T M N C K S  CAGTCTCCATTCTCCCTGACTGTGACAGCAGGAGAGAGGTCACTATGAATTGCAAGTCC
	G Q S L L N S V N Q R N Y L T W Y Q Q K GGTCAGAGTCTGTTAAACAGTGTAAATCAGAGGAACTACTTGACCTGGTACCAGCAGAAG
VL4715	PGQPPKLLINWASTRESGVPCCAGGGCAGCCTCCTAAACTGTTGATCTACTGGGCATCCACTAGGGAATCTGGAGTCCCT
	DOR F T A SOG S G T D F T L T I S S V Q  GATCGCTTCACAGCCAGTGGATCTGGAACAGATTTCACTCTCACCATCAGCAGTGTGCAG
L	A E D L A V Y Y C Q N D Y T Y P F T F G GCTGAAGACCTGGCAGTTATTACTGTCAGAATGATTATACTTATCCGTTCACGTTCGGA
Myc-tag	GGGGGGACCAAG <u>CTCGAG</u> ATCAAACGGGAACAAAAACTCATCTCAGAAGAGGGATCTGAAT
	TAATAAGATCAAACGGTAATAAGGATCCAGCTCGAATTC

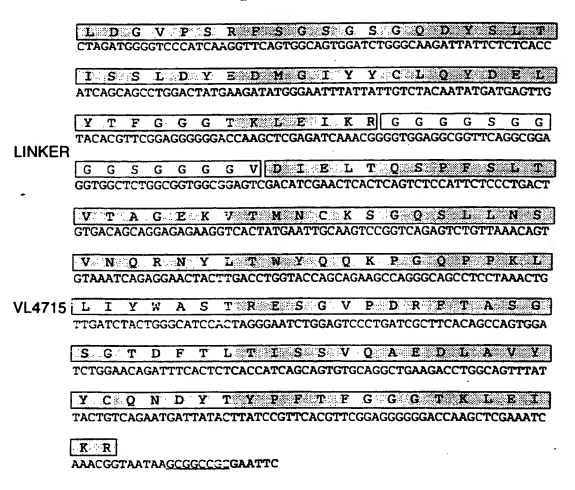
Fig.7. MK-YLLPT **AAGCTT**GCATGCAAATTCTATTTCAAGGAGACAGTCATAATGAAATACCTATTGCCTACG pelB LEADER AAGLLLLAAQPAMAQVQLQ GCAGCCGCTGGATTGTTATTACTCGCTGCCCAACCAGCGATGGCCCAGGTGCAGCTGCAG ESGGDLVKPGGSLTLSCATS GAGTCAGGGGGAGACTTAGTGAAGCCTGGAGGGTCCCTGACACTCTCTGTGCAACCTCT GOF T P 5 S Y AME S W V R O T S D K SEL GGATTCACTTTCAGTAGTTATGCCTTTTCTTGGGTCCGCCAGACCTCAGACAAGAGTCTG EWVATISSTEDTYTYYSDNVX VH4715 GAGTGGGTCGCAACCATCAGTAGTACTGATACTTATACCTATTATTCAGACAATGTGAAG GRFTISRDNGKNTLYLOM55 GGGCGCTTCACCATCTCCAGAGACAATGGCAAGAACACCCTGTACCTGCAAATGAGCAGT LKSEDTAVYYCARHGYYGKG CTGAAGTCTGAGGACACAGCCGTGTATTACTGTGCAAGACATGGGTACTATGGTAAAGGC Y F D Y W G Q G T T V T V S S G G G G S LINKER TATTTTGACTACTGGGGCCAAGGGACCACGGTCACCGTCTCCTCAGGTGGAGGCGGTTCA G G G G G G G S D I E L T Q S P F S L T V T A G E K V T M N C K S G O S L L CTGACTGTGACAGCAGGAGAGGAGGTCACTATGAATTGCAAGTCCGGTCAGAGTCTGTTA NSVRQRNYLTWYQQKPGQPP AACAGTGTAAATCAGAGGAACTACTTGACCTGGTACCAGCAGAAGCCAGGGCAGCCTCCT VL4715 K L L I Y W A S T R E S G V P D R F T A AAACTGTTGATCTACTGGGCATCCACTAGGGAATCTGGAGTCCCTGATCGCTTCACAGCC SGSGTDFTLTISSVQAEDLA AGTGGATCTGGAACAGATTTCACTCTCACCATCAGCAGTGTGCAGGCTGAAGACCTGGCA VYYCQNDYTYPFTFGGGTKL GTTTATTACTGTCAGAATGATTATACTTATCCGTTCACGTTCGGAGGGGGGGCCAAGCTC Myc-tag E I K R E Q K L I S E E D L N **GAGATCAAACGGGAACAAAAACTCATCTCAGAAGAGGATCTGAATTAATAAGATCAAACG GTAATAAGGATCCAGCTCGAATTC** 

Fig.8.1(2) MKYLLPT **AAGCTTGCATGCAAATTCTATTTCAAGGAGACAGTCATAATGAAATACCTATTGCCTACG** pelB AAAGLLLAAQPAMAQVQLQ LEADER GCAGCCCCTGGATTGTTATTACTCGCTGCCCAACCAGCGATGGCCCAGGTGCAGCTGCAG ESGGDLVSKIPGGSULTLSCATE GAGTCAGGGGGAGACTTAGTGAAGCCTGGAGGGTCCCTGACACTCTCCTGTGCAACCTCT VH4715 GFTPSSYAFSWVF078FKSL GGATTCACTTCAGTAGTTATGCCTTTTCTTGGGTCCGCCAGACCTCAGACAAGAGTCTG EWVATISMD DTYTYK BUN V E GAGTGGGTCGCAACCATCAGTAGTACTGATACTTATACCTATTATTCAGACAATGTGAAG GRFTLSRDNGKNTLYLQM55 GGGCGCTTCACCATCTCCAGAGACAATGGCAAGAACACCCTGTACCTGCAAATGAGCAGT LASEDITAVYYCARHGYTGK CTGAAGTCTGAGGACACAGCCGTGTATTACTGTGCAAGACATGGGTACTATGGTAAAGGC Y FDY W GOGTTVTVSSGGGGG TATTTTGACTACTGGGGCCAAGGGACCACGGTCACCGTCTCCTCAGGTGGAGGCGGTTCA G G G G G G G S A G S A Q V Q L Q Q LINKER GGCGGAGGTGGCTCTGGCGGTGGCGGATCGGCCGGTTCGGCCCAGGTCCAGCTGCAACAG S G P E L V R P G A S V K M S C K A S G TCAGGACCTGAGCTGGTAAAGCCTGGGGCTTCAGTGAAGATGTCCTGCAAGGCTTCTGGA TTFTSYVMHWVKQKPGGGLE TACACATTCACTAGCTATGTTATGCACTGGGTGAAACAGAAGCCTGGGCAGGGCCTTGAG WIGYIYPYNDGTKYNEKG TGGATTGGATATATTTATCCTTACAATGATGGTACTAAGTACAATGAGAAGTTCAAAGGC VH3418 KATLTSDKSSSTAYELLSSL **AAGGCCACACTGACTTCAGACAAATCCTCCAGCACAGCCTACATGGAGCTCAGCAGCCTG** TESEDS A VYVCSRAFDY WGQG ACCTCTGAGGACTCTGCGGTCTATTACTGTTCAAGACGCTTTGACTACTGGGGCCAAGGG TTVTVSS MKYLLPTAAAGLLLLA pelB LEADER GAGACAGTCATAATGAAATACCTATTGCCTACGGCAGCCGCTGGATTGTTATTACTCGCT AQPAMADIELTQSPSSMYAS GCCCAACCAGCGATGGCCGACATCGAGCTCACCCAGTCTCCATCTTCCATGTATGCATCT LGERITITCKASQDINTYLT CTAGGAGAGAATCACTATCACTTGCAAGGCGAGTCAGGACATTAATACCTATTTAACC VL3418

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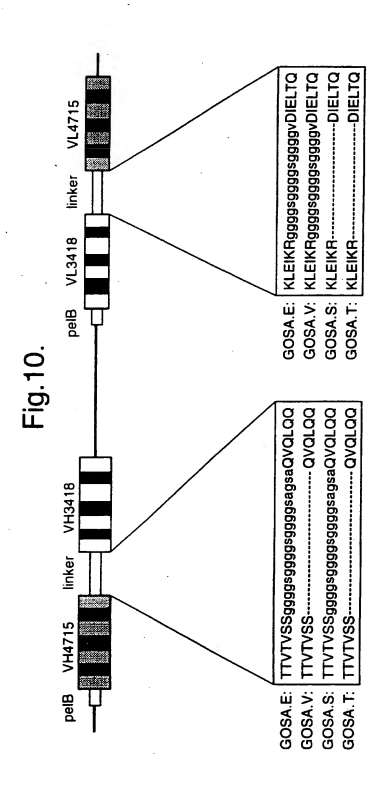
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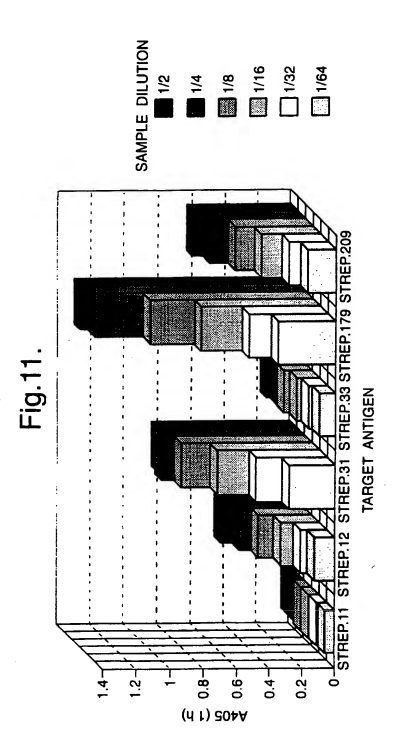
## Fig.8. 2(2)

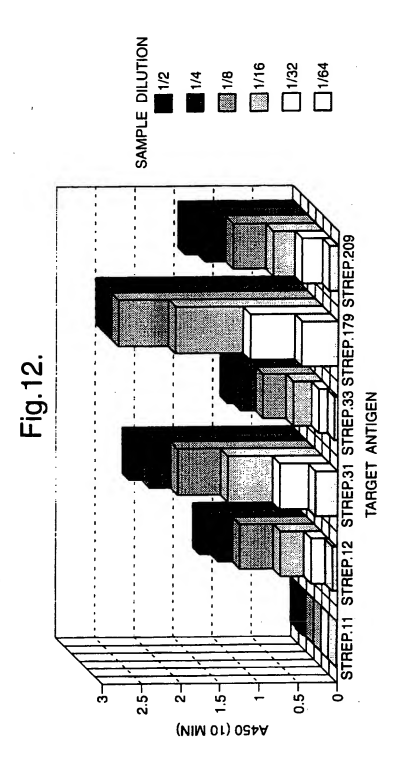




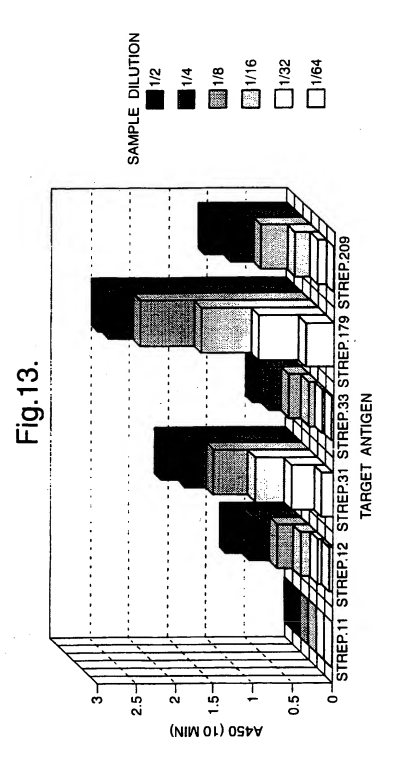
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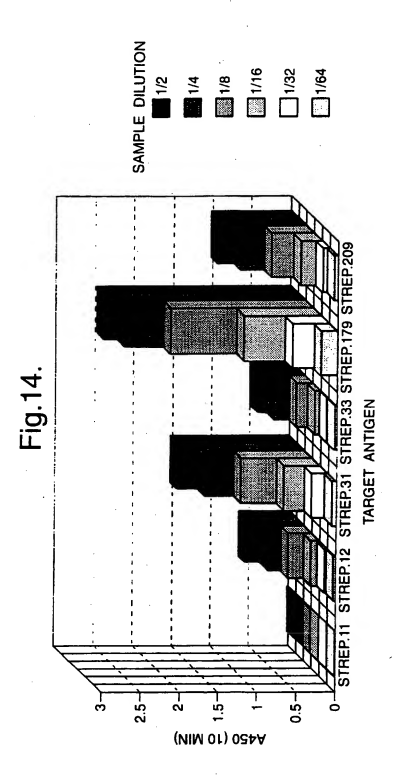


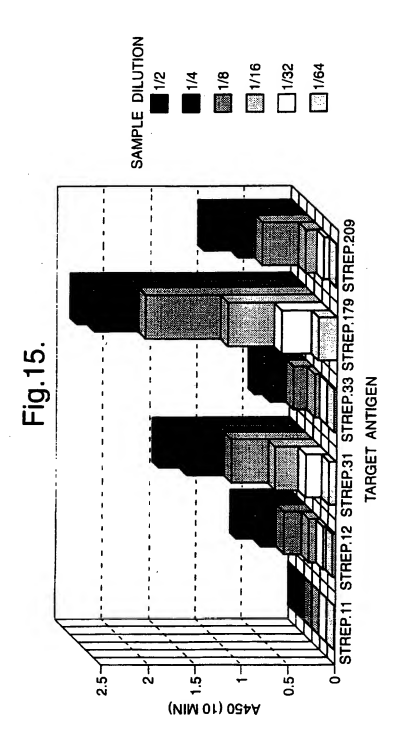




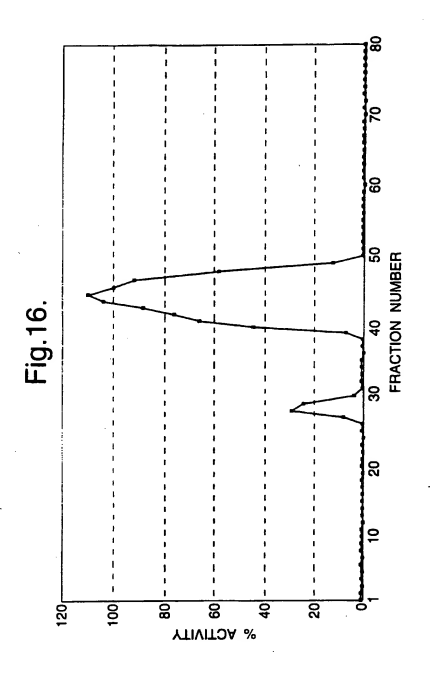
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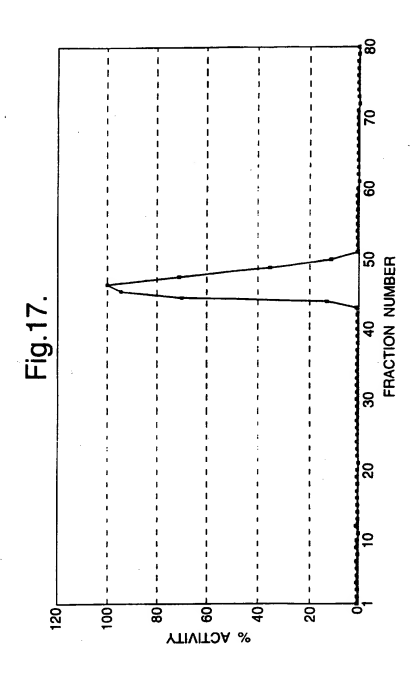






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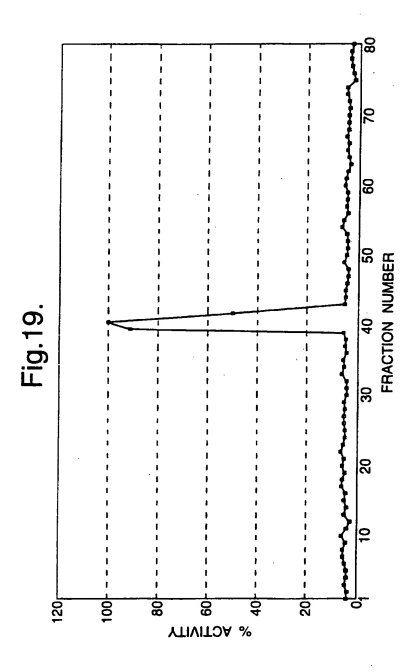
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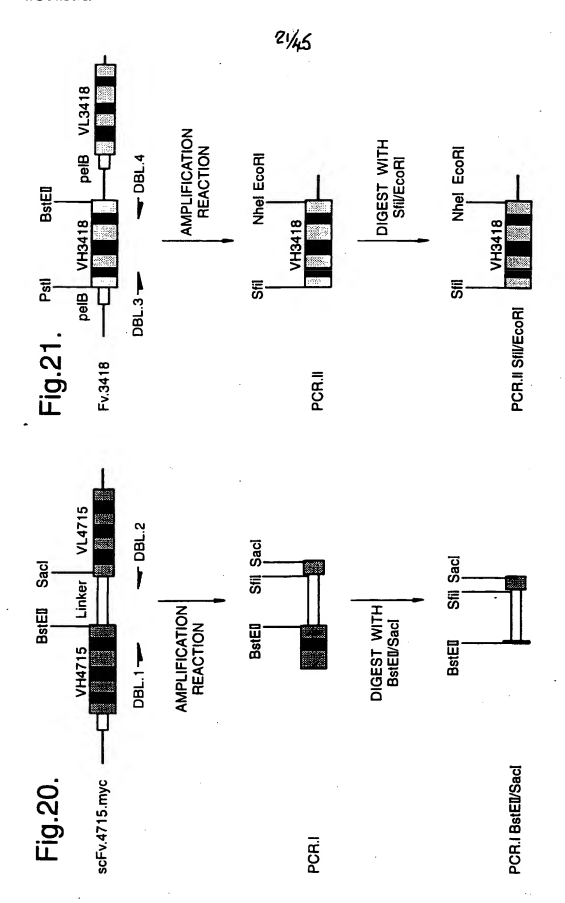
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FOR THE PURPOSES
OF INTERNATIONAL PROCESSING

(See Section 310(d)(ii) of the Administrative Instructions)

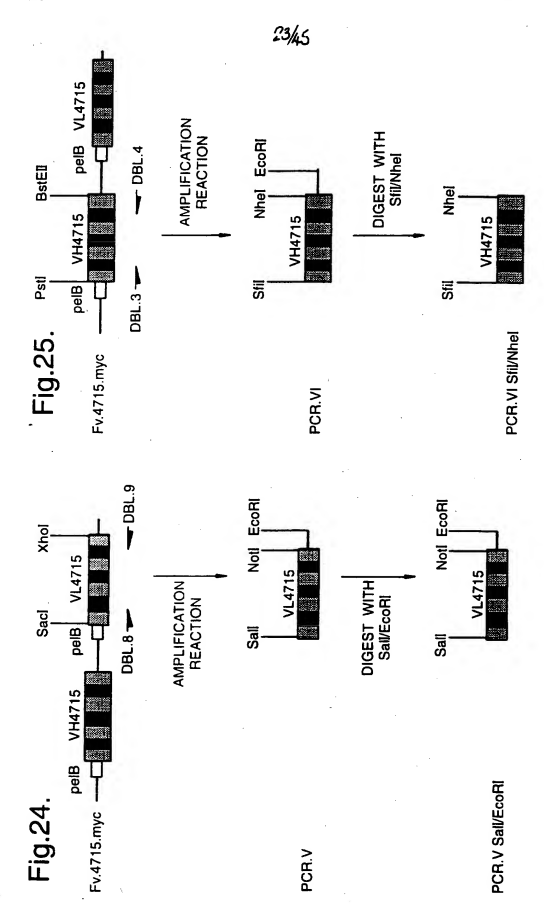


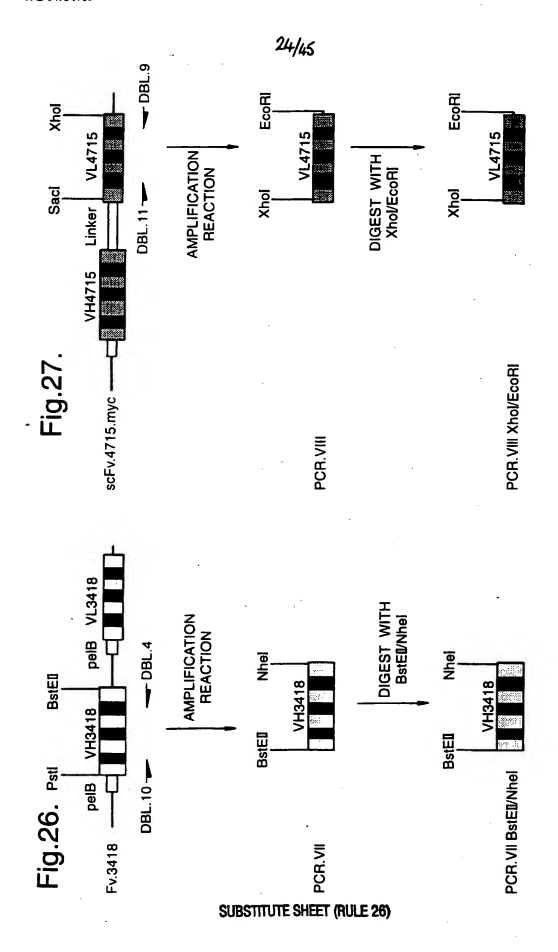


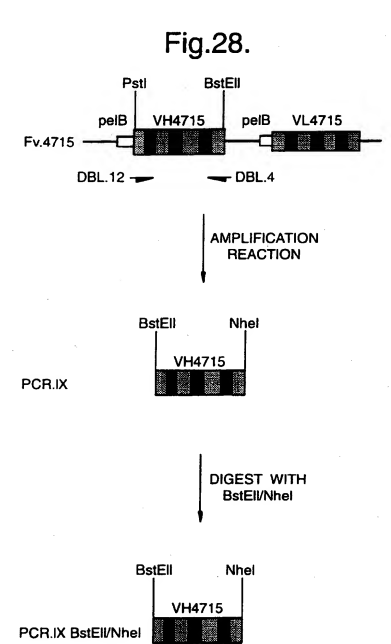
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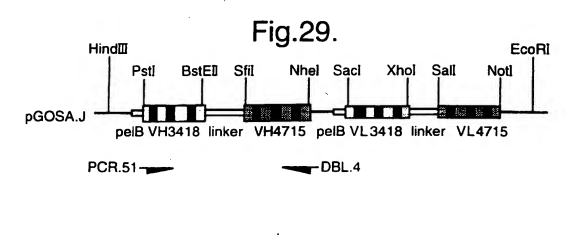


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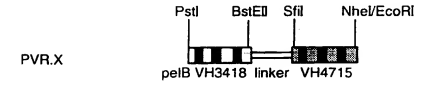








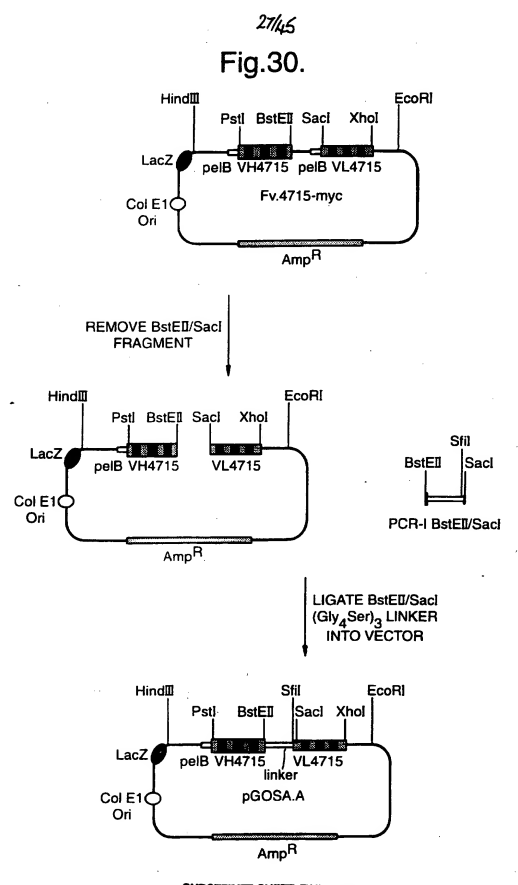
AMPLIFICATION REACTION





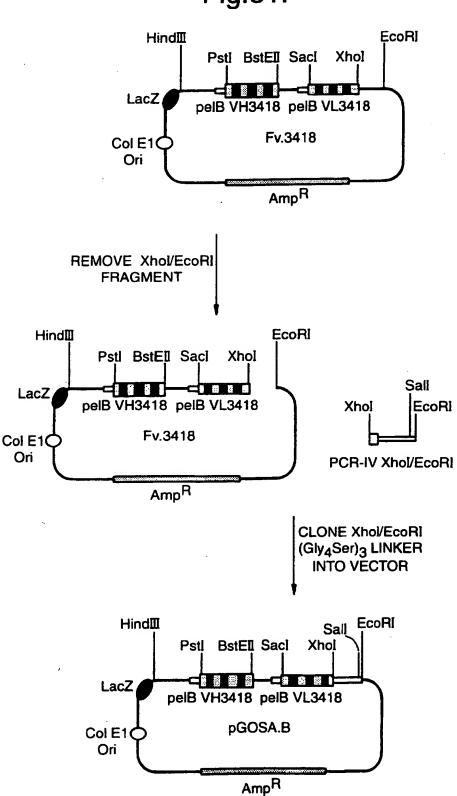


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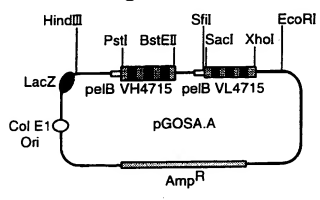
26/45 Fig.31.

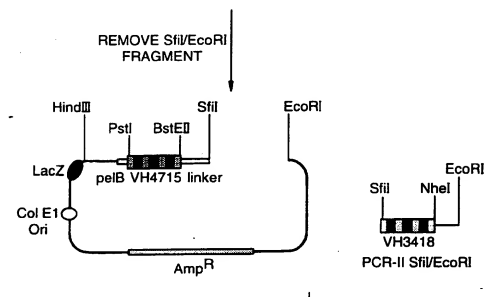


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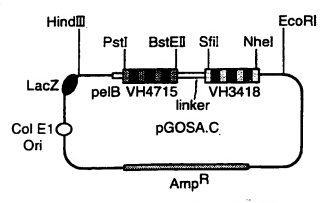


29/45 Fig.32.

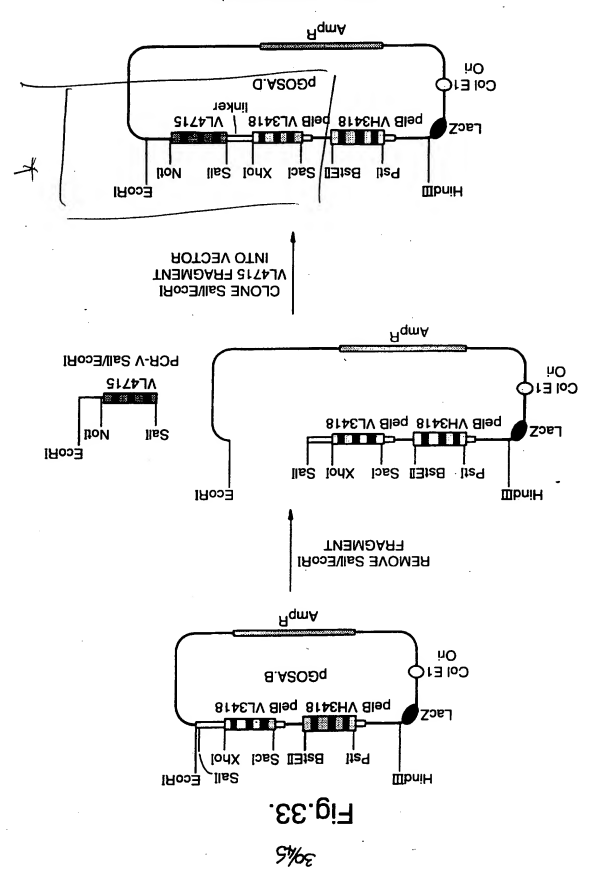




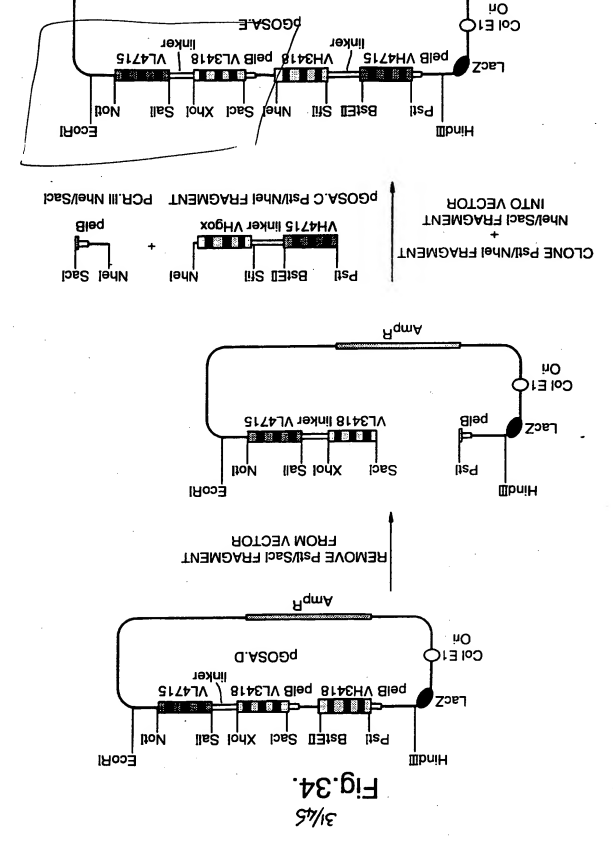




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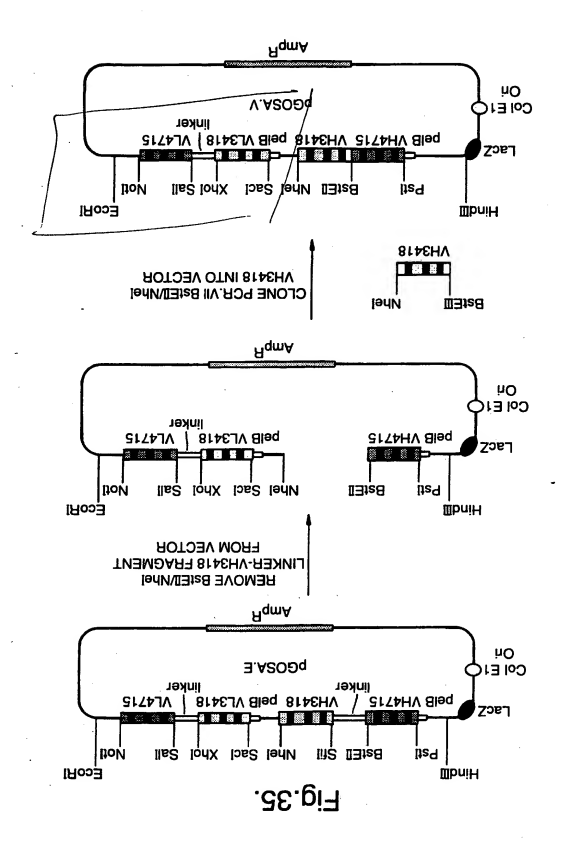
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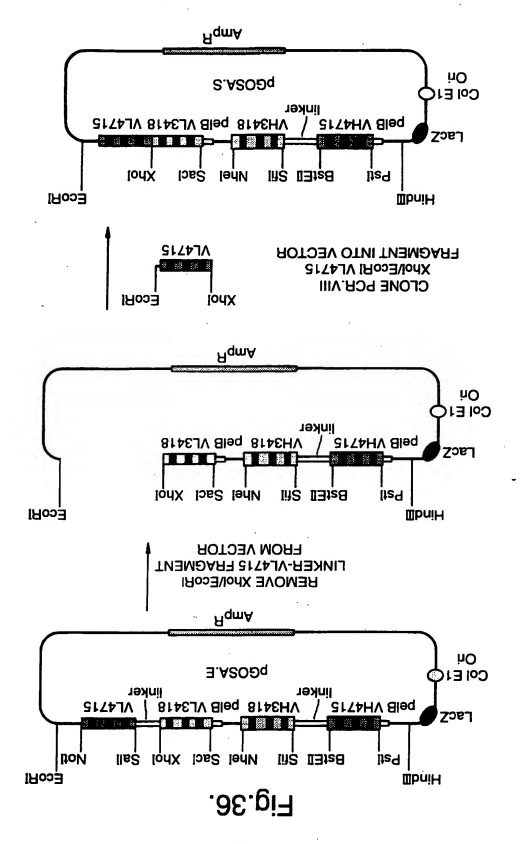


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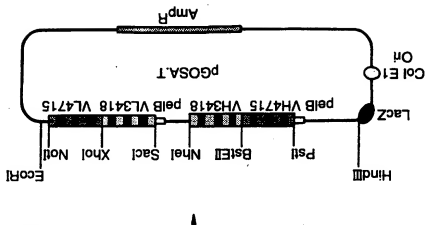
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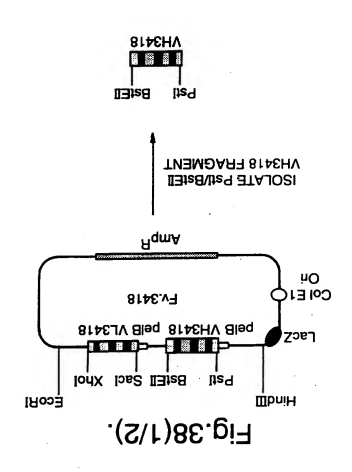


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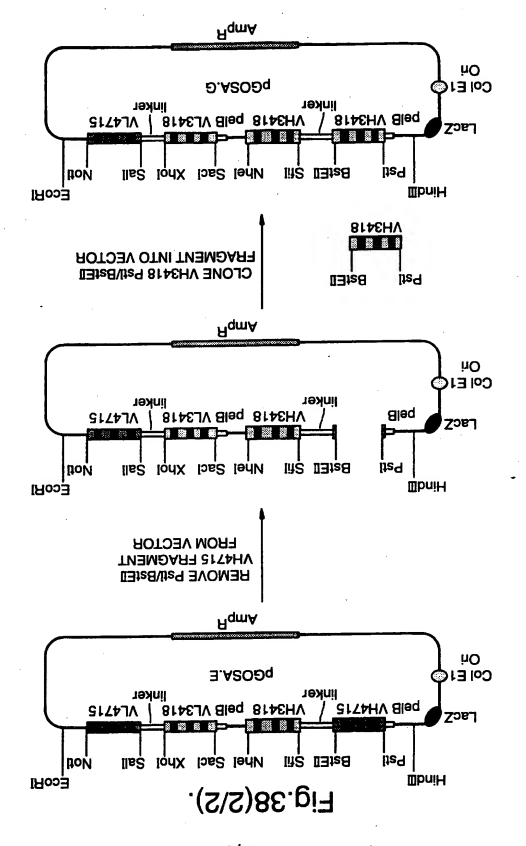
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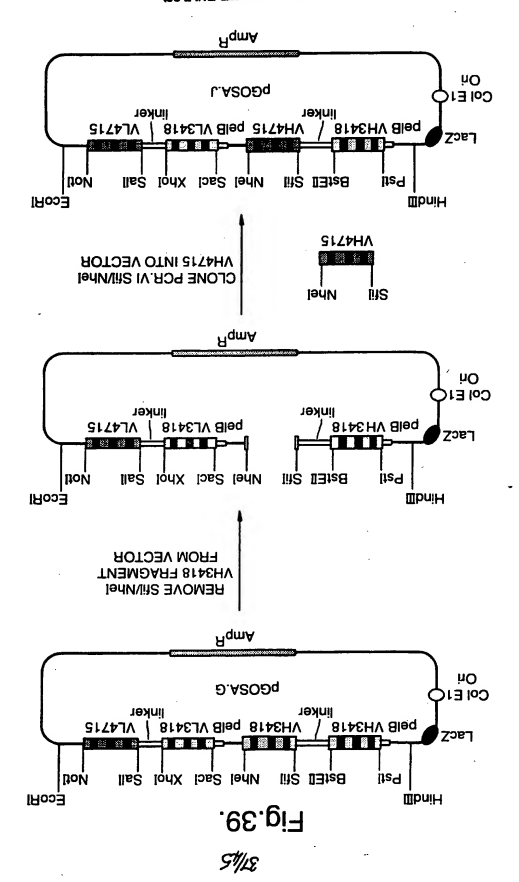
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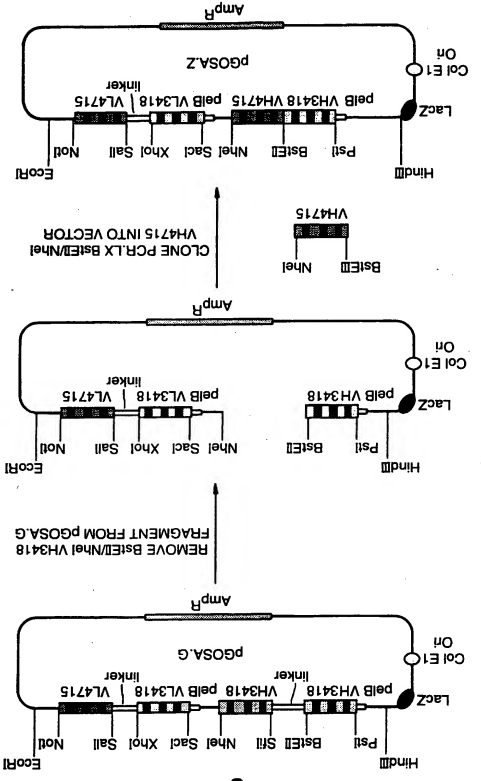


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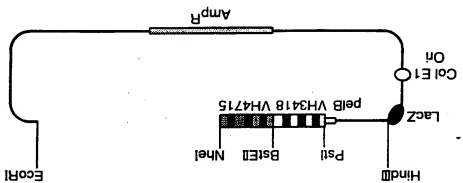
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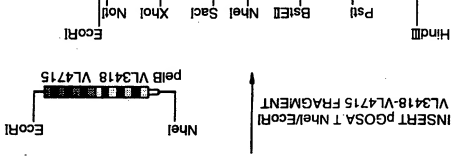
*3‰*5 Fig.40.

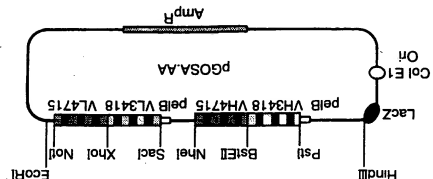


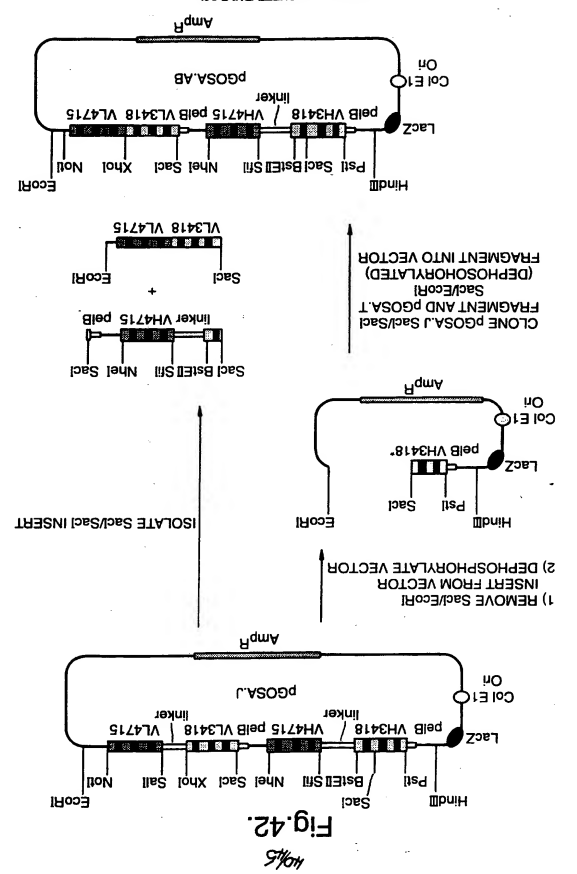
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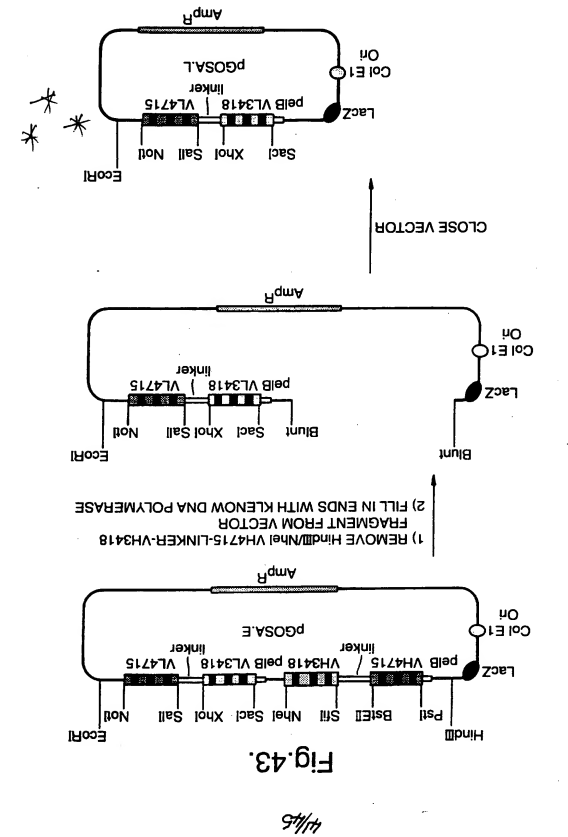






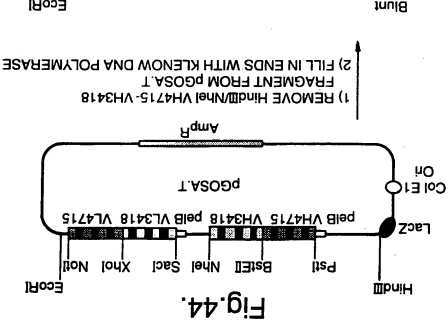


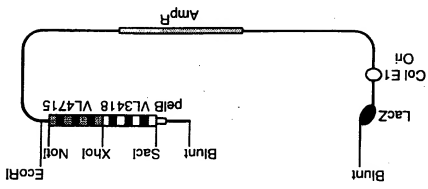
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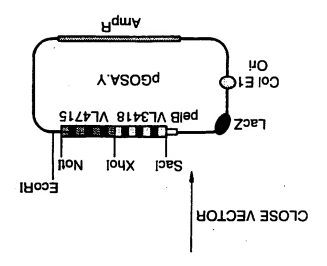


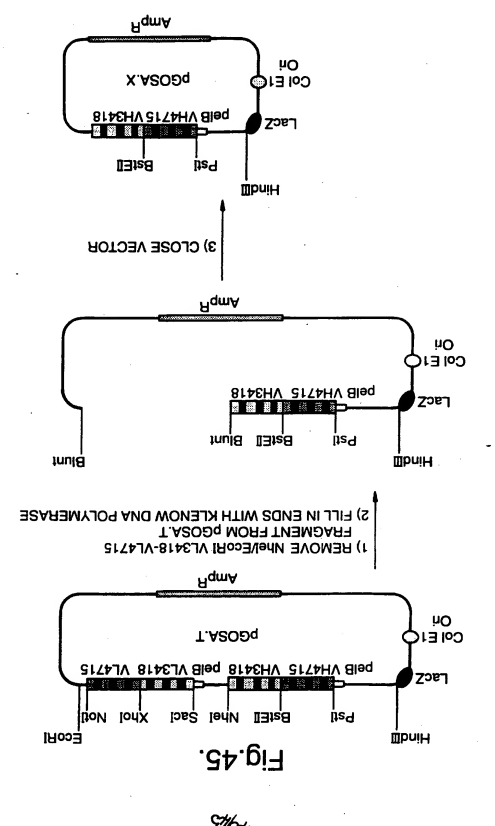
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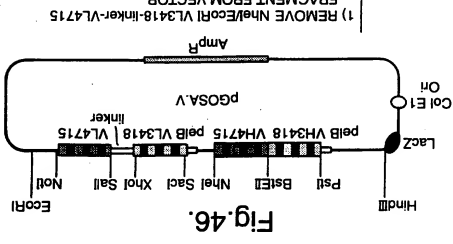




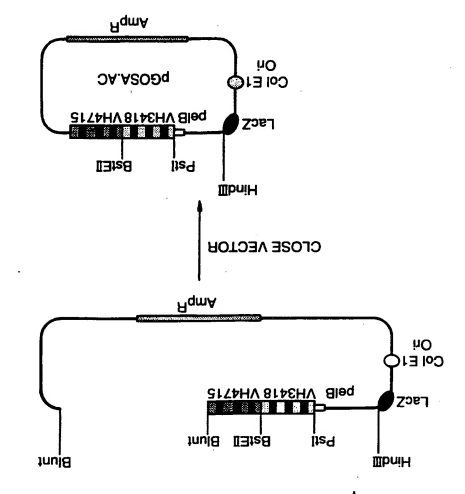


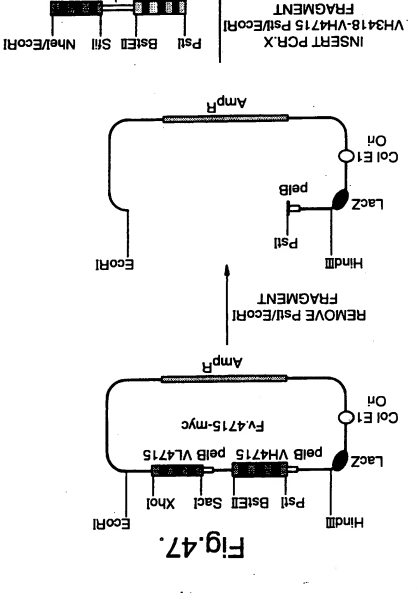
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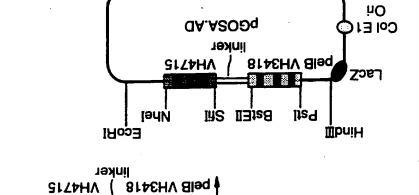




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